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## Commentary

# Q1 Curcumin as “Curecumin”: From kitchen to clinic

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### ABSTRACT

Although turmeric (*Curcuma longa*; an Indian spice) has been described in Ayurveda, as a treatment for inflammatory diseases and is referred by different names in different cultures, the active principle called curcumin or diferuloylmethane, a yellow pigment present in turmeric (curry powder) has been shown to exhibit numerous activities. Extensive research over the last half century has revealed several important functions of curcumin. It binds to a variety of proteins and inhibits the activity of various kinases. By modulating the activation of various transcription factors, curcumin regulates the expression of inflammatory enzymes, cytokines, adhesion molecules, and cell survival proteins. Curcumin also down-regulates cyclin D1, cyclin E and MDM2; and upregulates p21, p27, and p53. Various preclinical cell culture and animal studies suggest that curcumin has potential as an antiproliferative, anti-invasive, and antiangiogenic agent; as a mediator of chemoresistance and radioresistance; as a chemopreventive agent; and as a therapeutic agent in wound healing, diabetes, Alzheimer disease, Parkinson disease, cardiovascular disease, pulmonary disease, and arthritis. Pilot phase I clinical trials have shown curcumin to be safe even when consumed at a daily dose of 12 g for 3 months. Other clinical trials suggest a potential therapeutic role for curcumin in diseases such as familial adenomatous polyposis, inflammatory bowel disease, ulcerative colitis, colon cancer, pancreatic cancer, hypercholesterolemia, atherosclerosis, pancreatitis, psoriasis, chronic anterior uveitis and arthritis. Thus, curcumin, a spice once relegated to the kitchen shelf, has moved into the clinic and may prove to be “Curecumin”.

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## Q3 1. Introduction

16 Natural plant products have been used throughout human  
17 history for various purposes. Having coevolved with life, these  
18 natural products are billions of years old. Tens of thousands of  
19 them are produced as secondary metabolites by the higher  
20 plants as a natural defense against disease and infection.

21 Medicines derived from plants have played a pivotal role in the  
22 health care of many cultures, both ancient and modern [1–5].  
23 The Indian system of holistic medicine known as Ayurveda  
24 uses mainly plant-based drugs or formulations to treat various  
25 ailments including cancer. Of the approximately 877 small-  
26 molecule drugs introduced worldwide between 1981 and 2002,  
27 most (61%) can be traced back to their origins in natural  
28 products [1]. This is not surprising since plant-based drugs

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29 may be more suitable – at least in biochemical terms – for  
30 medicinal human use than the many exotic synthetic drugs  
31 produced through combinatorial chemistry. Nonetheless,  
32 modern medicine has neither held in very high esteem nor  
33 encouraged the medicinal use of natural products.

34 Over the last two decades, however, successful attempts to  
35 better understand molecular mechanisms of action of some  
36 natural products have kindled interest in their therapeutic use  
37 in modern medical settings. Remarkably, most of the natural  
38 products experimentally evaluated so far have been found to  
39 be nontoxic or to have effective doses far below their toxic  
40 doses. The role of natural products in human health care  
41 cannot be underestimated. An estimated 80% of individuals in  
42 developing countries depend primarily on natural products to  
43 meet their healthcare needs [6]. Recent surveys suggest that  
44 one in three Americans uses medicinal natural products daily  
45 and that possibly one in two cancer patients (i.e., up to 50% of  
46 patients treated in cancer centers) uses them as well. The  
47 current review is limited to curcumin, a natural product in use  
48 for thousands of years

49 Curcumin (diferuloylmethane), a polyphenol, is an active  
50 principle of the perennial herb *Curcuma longa* (commonly  
51 known as turmeric) (Fig. 1). The yellow-pigmented fraction of

52 turmeric contains curcuminoids, which are chemically related  
53 to its principal ingredient, curcumin. The major curcuminoids  
54 present in turmeric are demethoxycurcumin (curcumin II),  
55 bisdemethoxycurcumin (curcumin III), and the recently  
56 identified cyclocurcumin [7]. The major components of  
57 commercial curcumin are curcumin I (~77%), curcumin II  
58 (~17%), and curcumin III (~3%). The curcuminoid complex is  
59 also referred to as Indian saffron, yellow ginger, yellow root,  
60 *kacha haldi*, *ukon*, or natural yellow 3. Curcuminoids are  
61 present in 3–5% of turmeric. Though principally cultivated in  
62 India, Southeast Asia, China, and other Asian and tropical  
63 countries and regions, turmeric is also common in other parts  
64 of the world and is recognized by different names in different  
65 languages worldwide (Table 1) [8]

66 Curcumin was first isolated in 1815, obtained in crystalline  
67 form in 1870 [9,10], and ultimately identified as 1,6-hepta-  
68 diene-3,5-dione-1,7-bis(4-hydroxy-3-methoxyphenyl)-(1E,6E)  
69 or diferuloylmethane. In 1910, the feruloylmethane skeleton  
70 of curcumin was confirmed and synthesized by Lampe [11].  
71 Curcumin is a yellow-orange powder that is insoluble in water  
72 and ether but soluble in ethanol, dimethylsulfoxide, and  
73 acetone. Curcumin has a melting point of 183 °C, a molecular  
74 formula of C<sub>21</sub>H<sub>20</sub>O<sub>6</sub>, and a molecular weight of 368.37 g/mol.

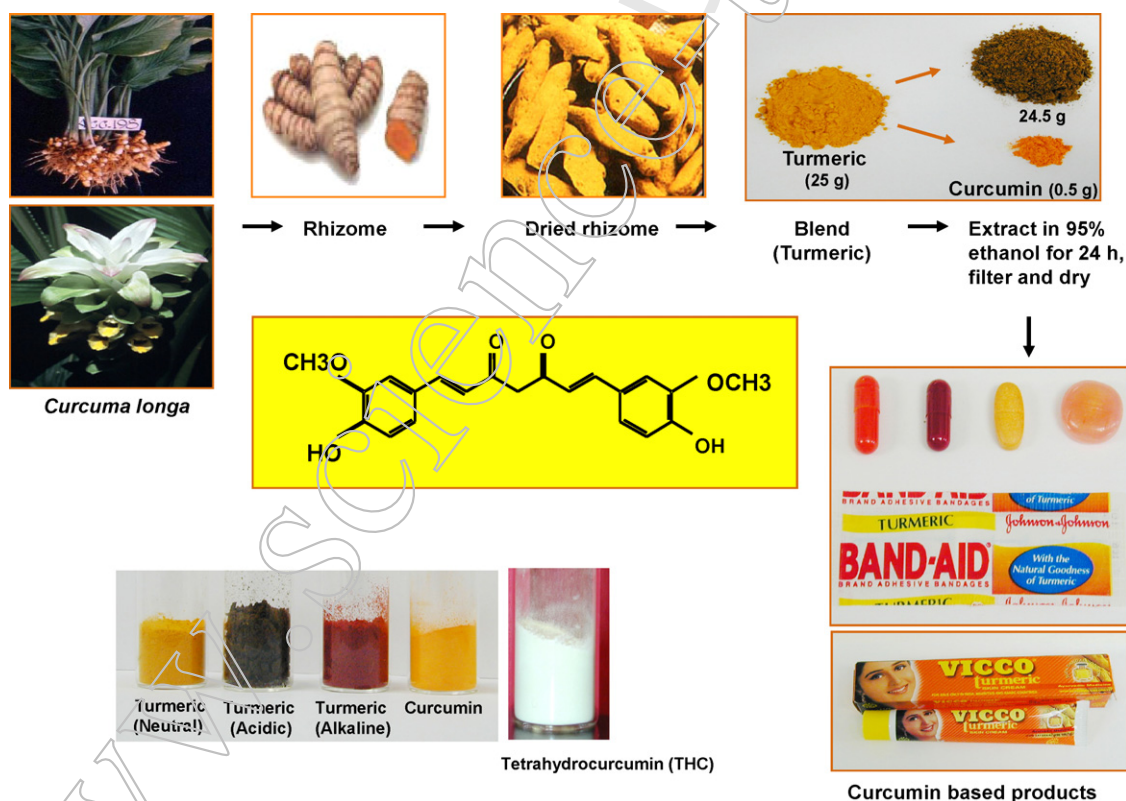


Fig. 1 – Isolation, extraction, and structure of curcumin. Curcumin capsules, pills, lozogens, band-aid and cream commonly sold in the market are shown. The change in color of turmeric at acidic and alkaline pH is also shown. Tetrahydrocurcumin (THC), a major metabolite of curcumin, exhibits whitish color. Alkaline turmeric (red color) is also referred as “Kumkum”. The traditional Kumkum, or Kungumam as it is called in Tamil Nadu (India), is made from dried turmeric. The turmeric is dried and powdered with a bit of slaked lime, which turns the rich yellow powder into red color. The kungumam (also called Bindi, Bindu, Tilak or Sandoor) is an auspicious symbol. When a girl or a married woman visits a house, it is a sign of respect (in case of an elderly lady) or blessings (in case of a young girl) to offer kumkum to them when they leave. Kumkum is also widely used for worshipping the Hindu goddesses, especially Shakti and Lakshmi. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of the article.)

Q16 **Table 1 – Various names of turmeric/curcumin in different languages**

| Language   | Name   |
|------------|--|
| Arabic     | Kurkum, Uqdah safra  |
| Armenian   | Toormerik, Turmerig  |
| Assamese   | Halodhi  |
| Bengali    | Halud  |
| Bulgarian  | Kurkuma  |
| Burmese    | Hsanwen, Sanwin, Sanae, Nanwin   |
| Catalan    | Cúrcuma  |
| Chinese    | Yu chin, Yu jin, Wohng geung, Geung wohng, Wat gam, Huang jiang, Jiang huang, Yu jin, Yu jin xiang gen                 |
| Croatian   | Indijski šafran, Kurkuma   |
| Czech      | Kurkuma, Indický Šafrán, Žlutý kořen, Žlutý zázvor   |
| Dhivehi    | Reen'dhoo  |
| Danish     | Gurkemeje  |
| Dutch      | Geelwortel, Kurkuma  |
| English    | Tarmeriek, Koenjit, Koenir   |
| Esperanto  | Indian saffron   |
| Estonian   | Kurkumo  |
| Farsi      | Harilik kurkuma, Kurkum, Pikk kollajuur, Lõhnnav kollajuur, Harilik kurkuma, Kurkum, Pikk kollajuur, Lõhnnav kollajuur |
| Finnish    | Zardchubeh   |
| French     | Kurkuma, Keltajuuri  |
| Galician   | Curcuma, Safran des Indes, Terre-mérite, Souchet des Indes   |
| German     | Cúrcuma  |
| Greek      | Curcuma, Kurkuma, Indischer Safran, Gelbwurz   |
| Gujarati   | Kitrinoriza, Kourkoumi, Kourkoumas   |
| Hebrew     | Halad, Haldar  |
| Hindi      | Kurkum   |
| Hungarian  | Haldi  |
| Hungarian  | Kurkuma, Sárga gyömbérgyökér   |
| Icelandic  | Túrmerik   |
| Indonesian | Kunyit, Kunir, Daun kunyit   |
| Italian    | Curcuma  |
| Japanese   | Ukon, Tamerikku  |
| Kannada    | Arishina, Arisina  |
| Khmer      | Romiet, Lomiet, Lamiet   |
| Korean     | Kang-hwang, Keolkuma Kolkuma, Sim-hwang, Teomerik, Tomerik, Tumerik, Ulgum, Uigumun                                    |
| Laotian    | Khi min khun, Khmin khun   |
| Latvian    | Kurkuma  |
| Lithuanian | Ciberžole, Kurkuma, Dažinė ciberžolė   |
| Malay      | Kunyit basah   |
| Malayalam  | Manjal   |
| Marathi    | Halad  |
| Nepali     | Haldi, Hardi, Besar  |
| Norwegian  | Gurkemeie  |
| Pahlavi    | Zard-choobag   |
| Pashto     | Zarchoba   |
| Polish     | Kurkuma, Ostryz' długi, Szafran indyjski   |
| Portuguese | Açafrão da Índia, Curcuma  |
| Punjabi    | Haldi  |
| Romanian   | Curcuma  |
| Russian    | Koren, kurkumy, Kurkuma  |

**Table 1 (Continued)**

| Language   | Name  |
|------------|---|
| Sanskrit   | Ameshta, bahula, bhadra, dhirgharaja, gandaplashika, gauri, gharshani, haldi, haridra, harita, hemaragi, hemaragini, hrivilasini, jayanti, jwarantika, kanchani, kaveri, krimighana, kshamada, kshapa, lakshmi, mangalaprada, mangalya, mehagni, nisha, nishaknya, nishawa, pavitra, pinga, pinja, pita, patavaluka, pitika, rabhanga, asa, ranjani, ratrimanika, shifa, shiva, shobhana, shyama, soughagouhaya, suvarna, suvarnavarna, tamasini, umavara, vauragi, varavarnini, varnadatri, varnini, vishagni, yamini, yohitapriya, yuvati |
| Singhalese | Kaha  |
| Slovak     | Kurkuma   |
| Slovenian  | Kurkuma   |
| Spanish    | Cúrcuma, Azafrán arabe  |
| Swahili    | Manjano   |
| Swedish    | Gurkmeja  |
| Tagalog    | Dilaw   |
| Tamil      | Manjal  |
| Telugu     | Haridra, Pasupu   |
| Thai       | Kha min chan, Kha min; Wanchakmadluk  |
| Tibetan    | Gaser, Sga ser  |
| Turkish    | Hint safrani, Sarı boya, Zerdeçal, Safran kökü, Zerdali, Zerdecöp, Zerdecube  |
| Ukrainian  | Kurkuma   |
| Urdu       | Haldi, Zard chub  |
| Vietnamese | Bot nghe, Cu nghe, Nghe, Uat kim, Khuong hoang  |
| Yiddish    | Kurkume   |

Modified from Ravindran et al. [8].

Spectrophotometrically, the maximum absorption ( $\lambda_{max}$ ) of curcumin in methanol occurs at 430 nm and in acetone at 415–420 nm [12]. A 1% solution of curcumin contains 1650 absorbance units. Curcumin appears brilliant yellow hue at pH 2.5–7 and red at pH > 7. Curcumin exists in enolic and  $\beta$ -diketonic forms. The fact that curcumin in solution exists primarily in its enolic form [13] has an important bearing on the radical-scavenging ability of curcumin.

The stability of curcumin in aqueous media improves at high pH (>11.7) [14,15]. Although quite soluble in organic solvents such as DMSO, ethanol, methanol, or acetone, it is poorly soluble in aqueous solvents [16]. Curcumin is stable at acidic pH but unstable at neutral and basic pH, under which conditions it is degraded to ferulic acid and feruloylmethane [15–17]. Most curcumin (>90%) is rapidly degraded within 30 min of placement in phosphate buffer systems of pH 7.2 [15,17]. The ability of antioxidants such as ascorbic acid, N-acetylcysteine (NAC), and glutathione to prevent this degradation suggests that an oxidative mechanism is at work. Degradation of curcumin is extremely slow at pH 1–6 [15], as normally encountered in the stomach. In contrast, one of curcumin's major metabolites (tetrahydrocurcumin, or THC) is quite stable at neutral or basic pH [18] and still possesses antioxidant activities [19–21]. Curcumin is soluble in 0.1 M sodium hydroxide, although it remains stable for only 1–2 h. In comparison, curcumin is more stable in cell culture medium

**Table 2 – A list of molecular targets of curcumin**

|  |
|--|
| Transcriptional factors                                  |
| Activating protein-1↓                                    |
| β-Catenin↓   |
| CREB-binding protein↓                                    |
| Early growth response gene-1↓                            |
| Electrophile response element↑                           |
| Hypoxia inducible factor-1↓                              |
| Notch-1↓   |
| Nuclear factor-kappa B↓                                  |
| Nuclear factor 2-related factor↑                         |
| Peroxisome proliferator-activated receptor-gamma↓        |
| Signal transducers and activators of transcription-1↓    |
| Signal transducers and activators of transcription-3↓    |
| Signal transducers and activators of transcription-4↓    |
| Signal transducers and activators of transcription-5↓    |
| Wilms' tumor gene 1↓                                     |
| Inflammatory cytokines                                   |
| Interleukin-1↓   |
| Interleukin-2↓   |
| Interleukin-5↓   |
| Interleukin-6↓   |
| Interleukin-8↓   |
| Interleukin-12↓  |
| Interleukin-18↓  |
| Monocyte chemoattractant protein↓                        |
| Migration inhibition protein↓                            |
| Macrophage inflammatory protein↓                         |
| Tumor necrosis factor alpha↓                             |
| Enzymes  |
| Arylamine N-acetyltransferases-1↓                        |
| ATFase↓  |
| ATPase↓  |
| Cyclooxygenase-2↓  |
| Desaturase↓  |
| DNA polymerase↓  |
| Farnesyl protein transferase↓                            |
| Gluthathione-S-transferase↑                              |
| Glutamyl cysteine ligase                                 |
| Hemeoxygenase-1↑   |
| Inducible nitric oxide synthase↓                         |
| Lipoxygenase↓  |
| Matrix metalloproteinase↓                                |
| NAD(P)H:quinone oxidoreductase↓                          |
| Ornithine decarboxylase↓                                 |
| Phospholipase D↓   |
| Src homology 2 domain-containing tyrosine phosphatase 2↑ |
| Telomerase↓  |
| Tissue inhibitor of metalloproteinase-3↓                 |
| Glutamate-cysteine ligase↑                               |
| Kinases  |
| Autophosphorylation-activated protein kinase↓            |
| Ca <sup>2+</sup> -dependent protein kinase↓              |
| EGF receptor-kinase↓                                     |
| Extracellular receptor kinase↓                           |
| Focal adhesion kinase↓                                   |
| IL-1 receptor-associated kinase↓                         |
| Janus kinase↓  |
| c-jun N-terminal kinase↑                                 |
| Mitogen-activated protein kinase↓                        |
| Phosphorylase kinase↓                                    |
| Protamine kinase↓  |
| Protein kinase A↓  |
| Protein kinase B↓  |
| Protein kinase C↓  |
| pp60c-src tyrosine kinase↓                               |
| Protein tyrosine kinase↓                                 |

**Table 2 (Continued)**

|  |
|--|
| Growth factors                             |
| Connective tissue growth factor↓           |
| Epidermal growth factor↓                   |
| Fibroblast growth factor↓                  |
| Hepatocyte growth factor↓                  |
| Nerve growth factor↓                       |
| Platelet derived growth factor↓            |
| Tissue factor↓                             |
| Transforming growth factor-β1↓             |
| Vascular endothelial growth factor↓        |
| Receptors                                  |
| Androgen receptor↓                         |
| Aryl hydrocarbon receptor↓                 |
| Chemokine (C-X-C motif) receptor 4↓        |
| Death receptor-5↑                          |
| EGF-receptor↓                              |
| Endothelial protein C-receptor↓            |
| Estrogen receptor-alpha↓                   |
| Fas receptor↑                              |
| Histamine (2)- receptor↓                   |
| Human epidermal growth factor receptor-2↓  |
| Interleukin 8-receptor↓                    |
| Inositol 1,4,5-triphosphate receptor↓      |
| Integrin receptor↓                         |
| Low density lipoprotein-receptor↓          |
| Adhesion molecules                         |
| Endothelial leukocyte adhesion molecule-1↓ |
| Intracellular adhesion molecule-1↓         |
| Vascular cell adhesion molecule-1↓         |
| Antiapoptotic proteins                     |
| B-cell lymphoma protein 2↓                 |
| Bcl-xL↓                                    |
| Inhibitory apoptosis protein-1 ↓           |
| Others                                     |
| Cyclin D1↓                                 |
| DNA fragmentation factor 40-kd subunit↑    |
| Heat-shock protein 70↑                     |
| Multi-drug resistance protein↓             |
| Urokinase-type plasminogen activator↓      |
| p <sup>53</sup> ↑                          |

For more information, see Ref. [43,44].

containing 10% fetal calf serum and in human blood, <20% of curcumin being degraded within 1 h and approximately 50% by 8 h [15]. *trans*-6-(4'-Hydroxy-3'-methoxyphenyl)-2,4-dioxo-5-hexenal is a major degradation product; vanillin, ferulic acid, feruloylmethane are minor degradation products. The amount of vanillin increases with incubation time. In addition, curcumin appears to be stabilized by forming complexes with cyclodextrin [22].

## 2. Traditional uses of curcumin

Traditionally, turmeric has been put to use as a foodstuff, cosmetic, and medicine. As a spice, it is used to provide curry with its distinctive yellow color and flavor. It is used as a coloring agent in cheese, butter, and other foods [23,24]. In folk medicine, turmeric and natural curcuminoids have been applied as therapeutic preparations over the centuries in different parts of the world. In Ayurvedic medicine, curcumin



is a well-documented treatment for various respiratory conditions (e.g., asthma, bronchial hyperactivity, and allergy) as well as for liver disorders, anorexia, rheumatism, diabetic wounds, runny nose, cough, and sinusitis [25]. In traditional Chinese medicine, it is used to treat diseases associated with abdominal pain [26]. In ancient Hindu medicine, it was used to treat sprains and swelling [25]. Throughout the Orient, it has traditionally been used to good therapeutic effect, particularly as an anti-inflammatory [12], and many of its therapeutic effects have been confirmed by modern scientific research. Such effects include antioxidant [27], anti-inflammatory [24,28,29], anticarcinogenic and antimicrobial [30–32], hepatoprotective [32], thrombosuppressive [33], cardiovascular (i.e., as protection against myocardial infarction) [29,34,35], hypoglycemic [36–38], and antiarthritic (i.e., as protection against rheumatoid arthritis) [39]. The most compelling and key rationale for the continuing traditional therapeutic use of curcumin is its extremely good safety profile. To date, no studies in either animals [40,41] or humans [42] have discovered any toxicity associated with the use of curcumin, and it is clear that curcumin is not toxic even at very high doses.

### 3. Molecular targets of curcumin

Accumulating evidence suggests that curcumin has a diverse range of molecular targets, which supports the notion that curcumin influences numerous biochemical and molecular cascades (Table 2). Among its molecular targets are transcription factors, growth factors and their receptors, cytokines, enzymes, and genes regulating cell proliferation and apoptosis.

#### 3.1. Curcumin interacts with numerous targets

Curcumin is apparently a highly pleiotropic molecule that interacts physically with its numerous targets (Table 3). It binds to and inhibits the activity of enzymes, growth factor receptors, metals, albumin, and other molecules. It binds proteins such as P-glycoprotein [68,69], multidrug resistance proteins 1 and 2 (MRP1 and MRP2) [59], glutathione [59], protein kinase C, ATPase [52,53], ErbB2 [61], and alpha1-acid glycoprotein (AGP) [50]. By directly binding small  $\beta$ -amyloid species, curcumin blocks aggregation and fibril formation in vitro and in vivo [51]. Curcumin irreversibly binds CD13/aminopeptidase N (APN) and inhibits tumor invasion and angiogenesis [55]. Curcumin has also been shown to inhibit the activity of lipoxygenase by binding lipoxygenase itself [65] or binding to phosphatidylcholine (PC) micelles and thereby inhibiting lipoxygenase 1 [74].

#### 3.2. Curcumin inhibits activation of transcription factors

Curcumin is a potent inhibitor of the activation of various transcription factors including nuclear factor- $\kappa$ B (NF- $\kappa$ B), activated protein-1 (AP-1), signal transducer and activator of transcription (STAT) proteins, peroxisome proliferator-activated receptor- $\gamma$  (PPAR- $\gamma$ ), and  $\beta$ -catenin [44]. These transcription factors regulate the expression of genes that contribute to

**Table 3 – Ligands that physically interact with curcumin**

|   |         |
|---|---------|
| Albumin   | [45–49] |
| Alfa-acid glycoprotein                            | [50]    |
| Amyloid protein                                   | [51]    |
| ATPase  | [52,53] |
| Autophosphorylation-activated protein kinase (AK) | [54]    |
| CD13/aminopeptidase N                             | [55]    |
| DNA polymerase-Y                                  | [56]    |
| Focal adhesion kinase                             | [57]    |
| Glutathione                                       | [58]    |
| GST-P1  | [60]    |
| HER2  | [61]    |
| Human alpha1-acid glycoprotein (AGP)              | [50]    |
| Iron, Cu <sup>2+</sup> , Zn <sup>2+</sup>         | [62,53] |
| Lipoxygenase                                      | [64,65] |
| Microtubulin                                      | [66]    |
| MRP 1 and 2                                       | [59]    |
| Nucleic acid                                      | [67]    |
| P-glycoprotein                                    | [68–70] |
| Phosphorylase kinase (Phk),                       | [54]    |
| Protein kinase A (PKA),                           | [54]    |
| Protein kinase C (PKC),                           | [54]    |
| Protamine kinase (cPK),                           | [54]    |
| pp60c-src tyrosine kinase                         | [54,57] |
| Thioredoxin reductase                             | [71]    |
| Topoisomerase II                                  | [72]    |
| Ubiquitin isopeptidase                            | [73]    |

tumorigenesis, inflammation, cell survival, cell proliferation, invasion, and angiogenesis.

#### 3.3. Curcumin downregulates the activity of multiple kinases

A variety of tyrosine kinases are activated by mutations that contribute to the malignant transformation, growth, and metastasis of human cancers. Accordingly, protein kinases involved in key growth signaling cascades are good candidate targets for novel chemopreventive approaches to treat many human cancers. For example, most human cancers over-express epidermal growth factor receptor (EGFR) and HER2/neu, which ultimately stimulates the proliferation of cancer cells [75]. Cellular experiments in vitro have shown that short-term treatment with curcumin inhibits EGFR kinase activity and EGF-induced tyrosine phosphorylation of EGFR in A431 cells and depletes cells of Her2/neu protein. Similar to geldanamycin, curcumin is extremely potent at degrading intracellular HER2 and disrupting its tyrosine kinase activity [76]. Additionally, as recently shown in our laboratory, curcumin may downregulate bcl-2 expression, thereby contributing to antiproliferative activity. Curcumin has also been shown to induce apoptosis in acute T cell leukemias by inhibiting the phosphatidylinositol 3 kinase/AKT pathway and to induce G2/M arrest and nonapoptotic autophagic cell death in malignant glioma cells by abrogating Akt and Erk signaling pathways [77].

Curcumin's effects are also apparently mediated through its inhibition of various other serine/threonine protein kinases. As we have previously shown, curcumin completely inhibits the activity of several protein kinases including phosphorylase kinase, protein kinase C (PKC), protamine kinase (cPK), autophosphorylation-activated protein kinase

(AK), pp60c-src tyrosine kinase. Other investigators have shown similar suppression of phorbol-12-myristate-13-acetate (PMA)-induced activation of cellular PKC by curcumin [43,44].

Most inflammatory stimuli typically activate 1 of 3 independent MAPK pathways leading to activation of the p44/42 MAPK (also called ERK1/ERK2), JNK, or p38 MAPK pathway, respectively. Curcumin can apparently inhibit all of these pathways directly or indirectly, thus providing evidence of its potent anti-inflammatory and anticarcinogenic effects [43,44].

### 3.4. Curcumin inhibits expression of growth and metastases promoting genes

Overexpression of oncogenes promotes tumor cell growth and provides an ideal platform on which to design chemopreventive regimens. Cyclooxygenase-2 (COX-2) is associated with a wide variety of cancers including cancers of the colon, lung and breast. Because of the importance of COX-2 inhibition in human carcinogenesis, much research in the past decade has been focused on the development of specific COX-2 inhibitors [78]. Several studies have shown that curcumin downregulates the expression of COX-2 protein in different tumor cell lines, most likely through the downregulation of NF- $\kappa$ B activation that is required for COX-2 activation. There is also evidence in the literature that curcumin-induced suppression of cell proliferation results in decreased cyclin D1 expression and CDK4-mediated retinoblastoma protein phosphorylation. As shown in hepatocellular cancer cells, curcumin appears to alter the metastatic potential of tumor cells by inhibiting the activity of matrix metalloproteinase-9 (MMP-9) and MMP-2 [79]. In experiments with ex vivo cultured BALB/c mouse peritoneal macrophages, curcumin reduced the production of iNOS mRNA in a concentration-dependent manner. Finally, curcumin appears to be able to exert anti-inflammatory and growth-inhibitory effects on cancer cells by inhibiting the expression of interleukin 1 $\beta$  (IL-1 $\beta$ ), interleukin 6 (IL-6), and tumor necrosis factor- $\alpha$  (TNF- $\alpha$ ) on the one hand and cyclin E on the other [80,81].

### 3.5. Curcumin inhibits expression of multiple genes/ pathways involved in apoptosis, cell invasion, and adhesion

Curcumin also operates through regulating the activities of additional molecular targets that control cell adhesion, apoptosis, and invasion. In this regard, curcumin has been shown to be an extremely potent inhibitor of TNF- $\alpha$ -induced expression of intracellular cell adhesion molecule-1 (ICAM-1), vascular cell adhesion molecule-1 (VCAM-1), and E-selectin in human umbilical vein endothelial cells. By apparently inhibiting the induction of steady-state transcription levels of ICAM-1, VCAM-1 and E-selectin, curcumin may be interfering detrimentally with the TNF- $\alpha$ -induced signaling event at an early stage. Additionally, curcumin has been shown to mediate its anticancer, chemosensitive, and radiosensitive effects via activation of p53 and simultaneous downregulation of MDM2 oncogene expression via the PI3K/mTOR/ETS2 pathway in human prostate cancer (PC3) and colon cancer (HT-29) cell lines [82,83] and to induce apoptosis and nuclear

translocation and activation of p53 in human neuroblastoma cells [84].

### 3.6. Curcumin regulates activities of several enzymes that mediate tumor growth

In addition to directly regulating the expression of candidate genes, curcumin also appears to effectively regulate the activities of enzymes that control tumor growth and proliferation. Curcumin blocks fibrosis in anti-Thy1 glomerulonephritis through its upregulation of hemoxygenase-1 (HO-1) gene expression, suggesting that it has antifibrotic effects in glomerular disease [85]. Similarly, curcumin can reportedly induce HO-1 expression through the generation of reactive oxygen species, p38 activation, and phosphatase inhibition [86].

Curcumin can also apparently suppress tumor cell growth through its effects on Ras protein pathways. Ras proteins, in order to extend their biological activity, must be isoprenylated at a conserved cysteine residue near the carboxyl terminus (Cys-186 in mammalian Ras p21 proteins). Previous studies have indicated that an intermediate in the mevalonate pathway, most likely farnesyl pyrophosphate, donates this isoprenyl group and that inhibitors of the mevalonate pathway might be able to block the transforming effects of Ras oncogenes expression. Indeed, in one study evaluating such a role for curcumin, curcumin derivatives strongly inhibited FPPase activity, thereby suggesting another potential mechanism by which curcumin might suppress cellular growth [43,44].

In another investigation, curcumin remarkably inhibited the activity of xanthine oxidase (XO) in vitro in PMA-treated NIH3T3 cells. Induction of XO activity is considered a major cause of PMA-mediated tumor promotion, and curcumin's marked ability to inhibit PMA-induced increases in such activity appears to lie in its direct inactivation of the XO protein [43,44].

## 4. Preclinical studies of curcumin

### 4.1. Curcumin is a potent chemopreventive agent

Numerous studies in rodent models argue for curcumin's chemopreventive potential in cancer (Table 4). Curcumin can reportedly suppress the tumorigenic activity of a wide variety of carcinogens in cancers of the colon, duodenum, esophagus, forestomach, stomach, liver, breast, leukemia, oral cavity, and prostate. In studies in mice, curcumin was able to inhibit 7,12-dimethylbenz[a]anthracene (DMBA)-initiated and 12-O-tetradecanoylphorbol-13-acetate (TPA)-promoted skin tumor formation [31,120,126]. Curcumin has also shown an ability to inhibit the mammary tumor-initiating activity of DMBA [110] and the in vivo formation of mammary DMBA-DNA adducts in female rats [111] and to exert chemopreventive activity when administered during the promotion/progression stage of colon carcinogenesis [91]. Meanwhile, one group has studied not only curcumin's chemopreventive effects but also its effects on the initiation or post-initiation phase of N-nitrosomethylbenzylamine (NMBA)-induced esophageal carcinogenesis in male F344 rats [100]. Using a slightly different approach,

**Table 4 – Curcumin exhibits chemopreventive effects against various cancers**

| Cancer                          | Carcinogen            | Animal         | Dose                    | Reference |
|---------------------------------|-----------------------|----------------|-------------------------|-----------|
| <b>Gastrointestinal cancers</b> |                       |                |                         |           |
| Aberrant crypt foci (ACF)       | Azoxymethane          | Rat            | 2000 ppm                | [87]      |
| Colon cancer                    | Azoxymethane          | Mice           | 0.5–0.2% (w/w)          | [88]      |
| Colon cancer                    | DMH                   | Mice           | 0.5%                    | [89]      |
| Colon cancer                    | Azoxymethane          | Rat            | 2000 ppm                | [90]      |
| Colon cancer                    | Azoxymethane          | Rat            | 0.2 or 0.6% (w/w)       | [91]      |
| Colon cancer                    | PhIP                  | Apc (min) mice | 2000 ppm                | [92]      |
| Colon cancer                    | Azoxymethane          | Rat            | 1 or 2% (w/w)           | [93]      |
| Colon cancer                    | Azoxymethane          | Rat            | 0.6% (w/w)              | [94]      |
| Colon cancer                    | 1,2-Dimethylhydrazine | Rat            | 0.6%                    | [95]      |
| Colitis                         | TNBS                  | Mice           | 0.5–5%, diet            | [96]      |
| Colitis                         | DNB                   | Mice           | 0.25%; diet             | [97]      |
| Colitis                         | TNBS                  | Mice           | 50 mg/kg                | [98]      |
| Ulcerative colitis              | DNCB                  | Rat            | 25–100 mg/kg            | [99]      |
| Duodenal tumor                  | MNNG                  | Mice           | 0.5–2.0% (w/w)          | [88]      |
| Esophageal cancer               | NMBA                  | Rat            | 500 ppm                 | [100]     |
| FAD                             | Azoxymethane          | Mice           | 2%                      | [101]     |
| FAP                             | –                     | Min/+ mice     | 0.1, 0.2 or 0.5% (w/w)  | [102]     |
| Forestomach neoplasia           | B[a]P                 | Mice           |                         | [103]     |
| Forestomach cancer              | B[a]P                 | Mice           | 2% (w/w)                | [104]     |
| Forestomach neoplasia           | B[a]P                 | Mice           |                         | [105]     |
| Stomach cancer                  | MNNG                  | Rat            | 0.05% (w/w)             | [106]     |
| <b>Liver cancers</b>            |                       |                |                         |           |
| Hepatic hyperplasia             | Diethylnitrosamine    | Rat            | 200 or 600 mg/kg        | [107]     |
| Liver cancer                    | Diethylnitrosamine    | Mice           | 0.2% (w/w)              | [107]     |
| <b>Lung cancers</b>             |                       |                |                         |           |
| Lung cancer                     | B[a]P and NNK         | A/J mice       | 2000 ppm                | [108]     |
| <b>Blood cancers</b>            |                       |                |                         |           |
| Lymphoma/leukemia               | DMBA                  | Sencar mice    | 2% (w/w)                | [109]     |
| <b>Breast cancers</b>           |                       |                |                         |           |
| Mammary tumor                   | DMBA                  | Rat            | 0.8–1.6% (w/w)          | [93]      |
| Mammary tumor                   | DMBA                  | Rat            | 50–200 mg/kg            | [110]     |
| Mammary tumor                   | DMBA                  | Rat            | 1% (w/w)                | [111]     |
| Mammary tumor                   | DMBA                  | Sencar mice    | 2% (w/w)                | [109]     |
| Mammary tumor                   | Gamma radiation       | Rat            |                         | [112]     |
| Mammary tumor                   | Gamma radiation       | Rat            | 1% (w/w)                | [113]     |
| Mammary tumor                   | DMBA                  | Rats           |                         | [114]     |
| Mammary tumor                   | DMBA                  | Sencar mice    |                         | [115]     |
| Mammary tumor                   | Gamma radiation       | Rat            |                         | [113]     |
| <b>Oral cancers</b>             |                       |                |                         |           |
| Oral cancer                     | MNA                   | Hamster        |                         | [116]     |
| Oral cancer                     | NQO                   | Rat            | 500 ppm                 | [117]     |
| <b>Prostate cancers</b>         |                       |                |                         |           |
| Prostate cancer                 | DMAB and PhIP         | Rat            | 15–500 ppm              | [118]     |
| <b>Skin cancers</b>             |                       |                |                         |           |
| Dermatitis                      | TPA + UV-A            | Mice           |                         | [119]     |
| Skin tumor                      | TPA                   | Mice           |                         | [120]     |
| Skin tumor                      | DMBA                  | Mice           |                         | [103]     |
| Skin tumors                     | TPA                   | Mice           | 10 and 30 $\mu$ mol     | [121]     |
| Skin tumor                      | TPA                   | Mice           |                         | [122]     |
| Skin tumor                      | TPA                   | Mice           | 1, 10, 100 or 3000 nmol | [123]     |
| Skin tumor                      |                       | Mice           |                         | [124]     |
| Skin tumor                      | DMBA                  | Mice           |                         | [105]     |
| Skin tumor                      | B[a]P and DMBA        | Mice           |                         | [101]     |
| <b>Other cancers</b>            |                       |                |                         |           |
| Multi-organ cancer              | DHPN, EHEN            | Rat            | 1% (w/w)                | [125]     |

Abbreviations: FAP, familial adenomatous polyposis; ACF, aberrant crypt foci; FAD, focal areas of dysplasia; B[a]P, benzo[a]pyrene; DMBA, 7,12-dimethylbenz[a]anthracene; TPA, 12-O-tetradecanoylphorbol-13-acetate; NNK, 4-(methyl-nitrosamino)-1-(3-pyridyl)-1-butanone; NQO, 4-nitroquinoline-1-oxidase; DMAB, 3,2'-dimethyl-4-aminobiphenol; PhIP, 2-amino-1-methylimidazo[4,5-b]pyridine; DHPN, 2,2'-dihydroxy-di-n-propylnitrosamine; EHEN, N-ethyl-N-hydroxyethylnitrosamine.

another group investigated curcumin's ability to prevent tumors in C57BL/6J-Min/+ (Min/+) mice that bear a germline mutation in the APC gene and spontaneously develop numerous intestinal adenomas by 15 weeks of age [127]. The data obtained in that study were corroborated by a later study of the effects of curcumin on apoptosis and tumorigenesis in male *apc* (min) mice treated with the human dietary carcinogen 2-amino 1-methyl-6-phenylimidazo[4,5-*b*]pyridine (PhIP) [92].

At least one study has examined curcumin's preventive effect on the development of adenomas in the intestinal tract of C57BL/6J-Min/+ mice, a model of human familial adenomatous polyposis (FAP) [102]. Another group reported that, during the initiation phases of azoxymethane-induced colonic carcinogenesis, azoxymethane inhibits the expression of colonic COX-1 expression without affecting that of COX-2 [128]. However, they also found that simultaneous treatment with dietary curcumin may increase COX-2 expression to compensate for the azoxymethane-induced reduction of COX-1 expression.

In another recent study, the effects of curcumin administered at a daily dose of 100 mg/kg were investigated in an animal (Wistar rat) model of *N*-nitrosodiethylamine (DENA)-initiated and phenobarbital (PB)-induced hepatocarcinogenesis [129]. In a recent follow-up study, the investigators in that study have substantiated this finding by reporting that 100 mg/kg curcumin daily prevented the reduction of defensive hepatic glutathione antioxidant activity, decreased lipid peroxidation, and minimized the histological alterations induced by DENA/PB [130]. In another study, investigators found that the administration of curcumin and a synthetic analog to nicotine-treated Wistar rats over a period of 22 weeks enhanced biochemical marker enzyme and lipid profiles [131]. In a study in rodents, curcumin was able to inhibit the development of *N*-methyl-*N'*-nitro-*N*-nitrosoguanidine (MNNG)-induced stomach cancer [106], an effect that

may be mediated in part by an ability to suppress the proliferation of *Helicobacter pylori* (the major pathogen in human gastric cancer) [132].

#### 4.2. Curcumin inhibits proliferation of tumor cells in vitro

Curcumin has the ability to inhibit the proliferation of an extremely wide array of cancer cell types in vitro. This includes cells from cancers of the bladder, breast, lung, pancreas, prostate, cervix, head and neck, ovary, kidney, and brain; and osteosarcoma, leukemia and melanoma [12].

#### 4.3. Curcumin exhibits antitumor activity in animals

Besides the extensive in vitro demonstrations of curcumin's antiproliferative effects, numerous other studies have evaluated its efficacy in various animal models in vivo (Table 5). The first animal studies of curcumin's antitumor effects – performed with ascitic lymphoma cells in mice – were reported in 1985 by Kuttan, et al. [133]. More recently, others have studied the antitumoral and inhibitory effects of curcumin on melanoma cells [141] and melanoma lung metastasis in mice [147].

Other studies in vivo have investigated the effects of curcumin on tumor angiogenesis and the biomarkers COX-2 and VEGF in hepatocellular carcinoma cells implanted in nude mice [148]. One group demonstrated that systemic administration of curcumin for 6 consecutive days to rats bearing the highly cachectic Yoshida AH-130 ascites hepatoma significantly inhibited tumor growth [149]. Meanwhile, others have shown that curcumin can suppress the growth of head and neck carcinoma [140], modulate the growth of prostate cancer in rodents [145], and inhibit the growth of human pancreatic cancer in nude mice, in part by suppressing angiogenesis and inducing apoptosis as reported recently [143].

**Table 5 – A list of studies describing antitumor effects of curcumin in animals**

| Tumor                 | Route    | Dose          | Model      | Reference |
|-----------------------|----------|---------------|------------|-----------|
| Ascites <sup>2</sup>  | i.p.     | 50 mg/kg      | Ascites    | [133]     |
| Ascites               | i.p.     | 50 mg/kg      | Ascites    | [134]     |
| Breast <sup>1</sup>   | Diet     | 2% (w/w)      | Orthotopic | [135]     |
| Breast <sup>1</sup>   | Diet     | 1% (w/w)      | Orthotopic | [136]     |
| Colon <sup>2</sup>    | i.v.     | 40 mg/kg      | Xenograft  | [137]     |
| Gastric cancer        | Oral     | 50–200 mg/kg  | Xenograft  | [138]     |
| Glioblastoma          | i.t.     | 10 mg/kg      | Orthotopic | [77]      |
| HCC <sup>3</sup>      |          | 100–200 mg/kg | Orthotopic | [139]     |
| Hepatoma              | Oral     | 50–200 mg/kg  | Xenograft  | [138]     |
| HNSCC <sup>4</sup>    | Sub cute | 50–250 μmol/L | Xenograft  | [140]     |
| Leukemia              | Oral     | 50–200 mg/kg  | Xenograft  | [138]     |
| Melanoma              | i.p.     | 25 mg/kg      | Xenograft  | [141]     |
| Ovarian               | i.p.     | 500 mg/kg     | Orthotopic | [142]     |
| Pancreas <sup>2</sup> | i.v.     | 40 mg/kg      | Xenograft  | [143]     |
| Pancreas              | Gavage   | 1 gm/kg       | Orthotopic | [144]     |
| Prostate              | Diet     | 2% (w/w)      | Xenograft  | [145]     |
| Prostate              | Gavage   | 5 mg/kg       | IV         | [146]     |
| Prostate              | Gavage   | 5 mg/day      | Xenograft  | [82]      |

1, Lung metastases; 2, liposomal curcumin; 3, intrahepatic metastasis; i.p., intraperitoneal; i.t., intratumoral; i.v., intravenous; HCC, hepatocellular carcinoma; HNSCC, head and neck squamous cell carcinoma.



380 More recent studies have evaluated curcumin's chemo-  
381 sensitizing and radiosensitizing effects. Our group [135]  
382 evaluated the chemosensitizing effect of curcumin in combi-  
383 nation with paclitaxel on breast cancer metastases to the lung.  
384 Others examined the effects of curcumin on human breast  
385 cancer (MDA-MB-231) cells in an immunodeficient mouse  
386 model of metastasis [136] and observed that the number of  
387 lung metastases significantly decreased after intercardiac  
388 injection of curcumin, a clear demonstration of curcumin's  
389 promise for dietary chemoprevention of metastases [136].

390 In our laboratory, we have recently investigated curcumin's  
391 effects alone and in combination against several cancers. We  
392 have found that (a) the combination of curcumin and  
393 gemcitabine inhibits pancreatic cancer growth in nude mice  
394 by inhibiting NF- $\kappa$ B regulated gene expression, cell prolifera-  
395 tion, and angiogenesis [144]; (b) the combination of curcumin  
396 and docetaxel is effective against human ovarian cancer in  
397 nude mice [142]; (c) curcumin can suppress the growth of  
398 human glioblastoma in rodents [77]; and (d) curcumin  
399 sensitizes colon cancers in nude mice to oxaliplatin [137]. In  
400 addition, other recent studies have shown that curcumin  
401 sensitizes prostate cancers to chemotherapeutics and radia-  
402 tion by downregulating expression of the MDM2 oncogene  
403 [82]. Together, these in vivo animal studies clearly suggest  
404 curcumin's anticancer potential when administered either  
405 alone or in combination with currently employed chemother-  
406 apeutic agents or radiation.

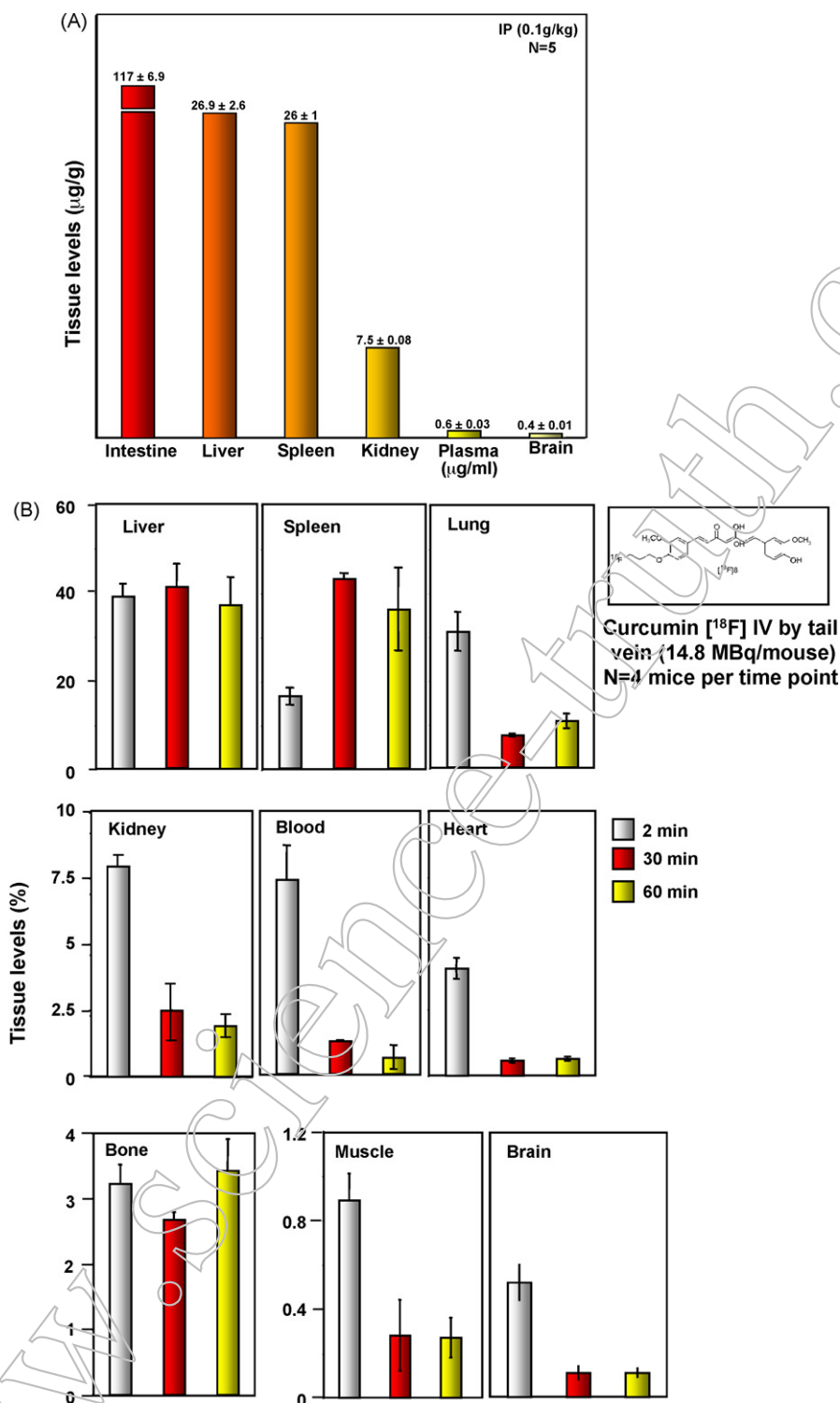
## 5. Pharmacokinetic and pharmacodynamic studies of curcumin in animals and humans

409 The pharmacokinetics and pharmacodynamics of curcumin  
410 have been widely investigated. Perhaps the first study to  
411 examine the uptake, distribution, and excretion of curcumin  
412 was conducted in 1978 by Wahlström and Blennow in  
413 Sprague-Dawley rats [150]. When administered orally at a  
414 dose of 1 g/kg, approximately 75% of the ingested curcumin  
415 was excreted in the feces and only negligible amounts in the  
416 urine. As indicated by blood plasma levels and biliary  
417 excretion, curcumin was poorly absorbed from the gut. No  
418 apparent toxic effects were seen after doses of up to 5 g/kg.  
419 When intravenously injected, curcumin was actively trans-  
420 ported into the bile. Most of the drug was metabolized,  
421 however, again suggesting poor absorption and rapid meta-  
422 bolism. Later, Holder et al. [151] administered deuterium- and  
423 tritium-labeled curcumin orally and intraperitoneally to rats  
424 and, like Wahlström and Blennow, found that most of it was  
425 excreted in the feces. When they administered curcumin  
426 intravenously and intraperitoneally to cannulated rats, the  
427 curcumin was excreted in the bile. The major biliary  
428 metabolites were glucuronides of tetrahydrocurcumin (THC)  
429 and hexahydrocurcumin (HHC); the minor biliary metabolite  
430 was dihydroferulic acid accompanied by traces of ferulic acid.  
431 In another study in which 400 mg curcumin was administered  
432 orally to rats, most of the administered curcumin (40%) was  
433 excreted unchanged in the feces, none in the urine (although  
434 curcumin glucuronide and sulfates were detected there), and  
435 none in heart blood (although traces were found in portal  
436 blood, liver, and kidney) [152]. Thirty minutes after adminis-

437 tration, 90% of the curcumin had appeared in the stomach and  
438 small intestine; by 24 h, only 1% remained there [152]. In  
439 another study by the same investigators, tritium-labeled  
440 curcumin administered at doses of 400, 80, and 10 mg was  
441 later detectable in the blood, liver, and kidney. At all three doses,  
442 the labeled curcumin was eliminated mainly through the feces  
443 and negligibly through the urine. At the two lowest doses (80  
444 and 10 mg), most of the labeled curcumin was excreted within  
445 72 h; conversely, at 400 mg, considerable amounts of labeled  
446 curcumin were still present in the tissues of interest 12 days  
447 after administration. The percentage of curcumin absorbed (60–  
448 66% of the given dose) remained constant regardless of the dose  
449 administered [153], indicating that increasing the dose of  
450 curcumin did not necessarily result in higher absorption.

451 In 1999, Pan et al. [18] investigated the pharmacokinetics of  
452 curcumin in mice. They found that, within the first 15 min  
453 after intraperitoneal (i.p.) administration of curcumin (0.1 g/  
454 kg), plasma curcumin levels had already reached 2.25  $\mu$ g/mL  
455 (Fig. 2). One hour after administration, curcumin levels in the Q5  
456 intestines, spleen, liver, and kidneys had reached 177.04,  
457 26.06, 26.90, and 7.51  $\mu$ g/g, respectively, but only trace levels  
458 (0.41  $\mu$ g/g) in the brain. In comparison, after oral administra-  
459 tion of 1 g/kg curcumin, serum plasma levels peaked at 0.5  $\mu$ M.  
460 Pan et al. also found curcumin-glucuronoside, dihydrocurcu-  
461 min-glucuronoside, THC-glucuronoside, and THC to be the  
462 major metabolites of curcumin in vivo. Together, these results  
463 agree with those of Ireson et al. [154,155], who examined  
464 curcumin metabolites in both rats and humans. As several  
465 groups have shown, the liver appears to be the major organ  
466 responsible for metabolism of curcumin [150,156,157]. Exam-  
467 ining rat liver tissue slices for the presence of curcumin  
468 metabolites, Hoehle and coworkers observed several reductive  
469 metabolites including THC, HHC, and octahydrocurcumin  
470 (OHC) and noted a predominance of OHC in males versus THC  
471 in females. They also identified both glucuronide and sulfate  
472 conjugates of THC, HHC, and OHC. This suggests that  
473 curcumin undergoes extensive reduction, most likely via  
474 alcohol dehydrogenase, before conjugation. In a Min/+ mouse  
475 model of FAP, Perkins et al. [102] examined the pharmaco-  
476 kinetics of curcumin administered either in the diet or in <sup>14</sup>C-  
477 labeled form as a single intraperitoneal dose. Though detected  
478 in only trace amounts in the plasma, curcumin was detected at  
479 levels ranging from 39 to 240 nmol/g in the small intestinal  
480 mucosa. The radiolabeled curcumin disappeared rapidly from  
481 tissues and plasma within 2–8 h after dosing. On the basis of  
482 their findings, Perkins et al. concluded that a daily dose of 1.6 g  
483 of curcumin is required for efficacy in humans. More recently,  
484 in a study examining the tissue distribution of radiolabeled  
485 fluoropropyl-substituted curcumin mice, Ryu et al. found that  
486 curcumin bound to  $\beta$ -amyloid plaques in the brain, thereby  
487 suggesting its possible use for brain imaging (Fig. 2) [158].

488 Pharmacokinetic studies in humans have generally pro-  
489 duced similar data though not always. In contrast to the case in  
490 rodents, oral dosing of curcumin at 4–8 g in one study resulted in  
491 peak plasma levels of 0.41–1.75  $\mu$ M [159]. In a small study of 15  
492 patients given oral curcumin (36–180 mg) daily for up to 4  
493 months, metabolites were not detected in the blood or urine but  
494 were detected in the feces [160]. In another study, Garcea et al.  
495 [161] examined the pharmacologically active levels of curcumin  
496 in patients with colorectal cancer who ingested curcumin at



**Fig. 2 - Plasma and tissue distribution of curcumin administered via intraperitoneal (i.p.) and systemic routes.** (A) Curcumin (0.1 g/kg) was administered (i.p.) to mice (N = 5), sacrificed 1 h later and concentration of curcumin in various tissues was analysed by HPLC. The data is replotted from [18]. (B) ICR mice were injected with [<sup>18</sup>F] labeled curcumin in 0.2 mL of 10% ethanol-saline via tail vein. The mice were sacrificed at the indicated times (2, 30, 60, and 120 min). Samples of blood, heart, lung, spleen, liver, spleen, kidney, muscle, brain, and bone were removed, weighed, and counted. Data are expressed as the percent injected dose per gram of tissue (% ID/g). The data is replotted from [158].

497 daily doses of 3600, 1800, or 450 mg for 7 days. By measuring  
498 curcumin's effects on the colorectal levels of DNA adduct 3-(2-  
499 deoxy- $\beta$ -di-erythro-penta-furanosyl)-pyr[1,2- $\alpha$ ]-purin-  
500 10(3H)one M(1)G and COX-2 protein, they showed that curcumin  
501 was taken up by both normal and malignant colorectal tissues  
502 and that it decreased M(1)G but not COX2 levels.

503 As most of these studies indicate, curcumin has poor  
504 bioavailability, and several groups have investigated ways to  
505 enhance it. Piperine has been shown to significantly enhance  
506 curcumin's bioavailability in studies involving both rats and  
507 healthy human volunteers. In brief, Shoba et al. [162]  
508 combined curcumin with piperine, a known inhibitor of  
509 hepatic and intestinal glucuronidation, and examined the  
510 resulting serum levels of curcumin. In the rat studies,  
511 administration of curcumin alone at a dose of 2 g/kg, resulted  
512 in moderate serum concentrations over 4 h. In contrast,  
513 concomitant administration with piperine 20 mg/kg increased  
514 for a short period the serum concentration of curcumin,  
515 significantly increased the time to maximum concentration  
516 while significantly decreasing elimination half-life and clear-  
517 ance, and increased bioavailability by 154%. In humans, on the  
518 other hand, administration of curcumin alone resulted in  
519 undetectable or trace amounts in the serum, whereas  
520 concomitant administration with piperine 20 mg/kg produced  
521 much higher concentrations and increased bioavailability by  
522 an astonishing 2000%. In another study in rats, other  
523 investigators found that a formulation of curcumin phosphatidylcholine given orally enhanced curcumin's bioavailability five-fold in plasma and in liver; but levels were lower in gastrointestinal mucosa [163]. Meanwhile, other attempts to increase the bioavailability of curcumin have been made, including the use of liposomal curcumin [143], nanoparticles of curcumin [164], and synthetic analogues of curcumin [165].

524 Whether curcumin metabolites are as active as curcumin  
525 itself is not clear. Although most studies indicate that  
526 curcumin glucuronides and THC are less active than curcumin  
527 [154,166], others suggest otherwise [20,21,89,167-172]. The  
528 differences in results so far are most likely due to the assays  
529 employed. For example, the phenolic glucuronides of curcu-  
530 min and its natural congeners, but not the parent compounds,  
531 have been shown to inhibit the assembly of microtubule  
532 proteins under cell-free conditions, implying that the glucur-  
533 onides are chemically reactive [167].

## 540 6. Clinical studies of curcumin

541 In response to the growing mass of in vitro and in vivo  
542 evidence for curcumin's chemopreventive and therapeutic  
543 efficacy, a number of clinical trials over the past two and a half  
544 decades have addressed the pharmacokinetics, safety, and  
545 efficacy of curcumin in humans (Table 6). Although these trials  
546 have concerned numerous inflammatory diseases including  
547 cancer, our focus in the sections to come will be on those  
548 dealing with cancers.

### 549 6.1. Curcumin is extremely safe and well tolerated

550 The potential use of curcumin in chemopreventive or  
551 therapeutic settings has raised the obvious issues of toxicity

and tolerance. At least three different phase I clinical trials  
552 indicate that curcumin is well tolerated when taken at doses  
553 as high as 12 g/day [159,162] (Table 6). These results were  
554 recently confirmed in an elegant dose-escalation trial to  
555 determine curcumin's maximum tolerated dose and safety  
556 [193]. In that trial, a standardized powder extract of uniformly  
557 milled curcumin (C3 Complex™, Sabinsa Corporation), was  
558 administered to 24 healthy volunteers at single doses ranging  
559 from 500 to 12,000 mg. Remarkably, only minimal, non-dose-  
560 related toxicity was seen and then only in seven subjects  
561 (30%). No curcumin was detected in the serum of subjects  
562 administered 500, 1000, 2000, 4000, 6000 or 8000 mg and only  
563 low levels in two subjects administered 10,000 or 12,000 mg.  
564

### 565 6.2. Curcumin has anti-inflammatory and antirheumatic activity

566 Rheumatoid arthritis is a frequent complication in the elderly,  
567 and most treatments aim at reducing the temporary symp-  
568 toms attributable to the underlying inflammatory activity  
569 [194]. The need for new treatment approaches has led to the  
570 recent introduction of potent disease-modifying antirheu-  
571 matic drugs (DMARDs), whose clinical benefits are unfortu-  
572 nately offset by their high cost and frequently undesirable side  
573 effects. Curcumin has been considered as an alternative.  
574

575 In the first clinical trial of curcumin's efficacy as an  
576 antirheumatic, investigators compared its antirheumatic  
577 potential with that of phenylbutazone in a short-term,  
578 double-blind, crossover study involving 18 relatively young  
579 patients (age range, 22-48 years) [39]. Each subject received a  
580 daily dose of either curcumin (1200 mg) or phenylbutazone  
581 (300 mg) for 2 weeks. At the dose used, curcumin was well  
582 tolerated, had no side effects, and exerted an antirheumatic  
583 activity comparable to that of phenylbutazone.

584 Meanwhile, in a study of curcumin's anti-inflammatory  
585 properties, Satoskar et al. [173] evaluated curcumin's effects  
586 on spermatic cord edema and tenderness in 46 men between  
587 15 and 68 years old who had just undergone surgical repair of  
588 an inguinal hernia and/or hydrocele. After surgery, subjects  
589 were randomly assigned to receive curcumin (400 mg),  
590 phenylbutazone (100 mg), or placebo (250 mg lactose) three  
591 times a day on postoperative days 1-5. As in a previous study  
592 by Deodhar et al. [39], curcumin was deemed quite safe and,  
593 along with phenylbutazone, elicited much better anti-inflam-  
594 matory responses than placebo did [173].

### 595 6.3. Curcumin has potential as palliative therapy for cancerous skin lesions

596 External sebaceous neoplasms (e.g., actinic keratosis, super-  
597 ficial basal cell carcinoma, and external genital warts) have  
598 traditionally been treated topically with corticosteroid creams.  
599 In a study by Kuttan et al. [174], curcumin's efficacy when  
600 applied as either an ethanol extract of turmeric or as an  
601 ointment to external cancerous skin lesions was evaluated in  
602 62 patients. Regardless of the application, curcumin provided  
603 remarkable symptomatic relief that was in many cases  
604 relatively durable (lasting several months) and in all cases  
605 (except for a single adverse reaction in one subject) extremely  
606 safe. Its effects included less itching in almost all cases,  
607

**Table 6 – A list of clinical trials with curcumin in patients with different diseases**

| Disease  | Dose/frequency                           | Patients        | End point modulation  | Reference |
|--|--|-----------------|---|-----------|
| <b>Safety trials</b>                                   |  |                 |   |           |
| Phase 1  | 2000 mg/day <sup>1</sup>                 | 10              | Piperine enhanced bioavailability by 2000%  | [162]     |
| Phase-I  | 500–12,000 mg/day × 90 days              | 25              | Histologic improvement of precancerous lesions <sup>4</sup>   | [159]     |
| Phase 1  | 500–12,000 mg/day                        | 24              | Safe, well-tolerated even at 12 g/day   | [42]      |
| <b>Efficacy trials</b>                                 |  |                 |   |           |
| Rheumatoid arthritis                                   | 1200 mg/day × 14 days                    | 18              | Improved symptoms   | [39]      |
| Postoperative inflammation                             | 400 mg; 3×/day × 5 days                  | 46              | Decrease in inflammation  | [173]     |
| External cancerous lesions                             | 1% ointment × several months             | 62              | Reduction in smell in 90% patients, reduction of itching in all cases, dry lesions in 70% patients, reduction in lesion size and pain in 10% patients | [174]     |
| Cardiovascular   | 500 mg/day × 7 days                      | 10              | Decreased serum lipid peroxidase (33%), increased HDL cholesterol (29%), decreased total serum cholesterol (12%)                                      | [175]     |
| Atherosclerosis  | 10 mg; 2×/day × 28 days                  | 12              | Lowered LDL and apoB, increased HDL and ApoA  | [176]     |
| HIV  | 625 mg; 4×/day × 56 days                 | 40              | Well tolerated  | [177]     |
| Gall bladder function                                  | 20 mg, single dose (2 h)                 | 12              | Decreased gall bladder volume by 29%  | [178]     |
| Gall bladder function                                  | 20–80 mg, single dose (2 h)              | 12              | Decreased gall bladder volume by 72%  | [179]     |
| Chronic anterior uveitis                               | 375 mg; 3×/day × 84 days                 | 32              | Eighty-six percent decrease in chronic anterior uveitis   | [180]     |
| Idiopathic Inflammatory Orbital Pseudotumors           | 375 mg; 3×/day × 180–660 days            | 8               | Four patients recovered completely. One patient showed decrease in swelling, no recurrence  | [181]     |
| Psoriasis  | 1% curcumin gel                          | 40              | Decreased PhK <sup>2</sup> , TRR <sup>3</sup> , parakeratosis, and density of epidermal CD8+ T cells  | [182]     |
| Colorectal cancer                                      | 36–180 mg/day × 120 days                 | 15              | Lowered GST   | [160]     |
| Colorectal cancer                                      | 450–3600 mg/day × 120 days               | 15              | Lowered inducible serum PGE2 levels   | [183]     |
| Irritable bowel syndrome                               | 72–144 mg/day × 56 days                  | 20 <sup>7</sup> | Reduced symptoms  | [184]     |
| Liver metastasis of CRC                                | 450–3600 mg/day × 7 days                 | 12              | Low bioavailability   | [156]     |
| Colorectal cancer                                      | 450–3600 mg/day × 7 days                 | 12              | Decreased M1G DNA adducts   | [161]     |
| Cadaveric renal transplantation                        | 480 mg; ×1–2/day × 30 days               | 43              | Improved renal function, reduced neurotoxicity  | [185]     |
| Tropical pancreatitis                                  | 500 mg/day × 42 days                     | 20              | Reduction in the erythrocyte MDA levels. Increased in erythrocyte GSH levels  | [186]     |
| Ulcerative proctitis                                   | 550 mg; × 2–3/day × 60 days              | 5               | Improved symptoms   | [187]     |
| Crohn's disease  | 360 mg; ×3/day × 30 days; ×4 for 60 days | 5               | Improved symptoms   | [187]     |
| Ulcerative colitis                                     | 2000 mg/day × 180 days                   | 89              | Low recurrence; improved symptoms   | [188]     |
| Familial adenomatous polyposis                         | 480 mg; ×3/day × 180 days                | 5               | Decrease in the number of polyps was 60.4%. Decrease in the size of polyps was 50.9%  | [189]     |
| Improves cognitive function                            | –  | 1010            | Better MMSE score <sup>5</sup>  | [190]     |
| Prostatic intraepithelial neoplasia (PIN) <sup>1</sup> | –  | 24              | –   | [191]     |
| <i>Helicobacter pylori</i> infection <sup>2</sup>      | 300 mg/day × 7 days                      | 25              | Significant improvement of dyspeptic symptoms   | [192]     |

Note: 1, + piperine 20 mg/kg; 2, PhK: phosphorylase kinase; 3, TRR: keratinocyte transferrin receptor; 4, histologic improvement of precancerous lesions was seen in one out of two patients with recently resected bladder cancer, two out of seven patients of oral leucoplakia, one out of six patients of intestinal metaplasia of the stomach, one out of four patients with CIN and two out of six patients with Bowen's disease; 5, MMSE: Mini-Mental State Examination Score; 1, Zyflamend, a polyherbal preparation containing curcumin was used; PIN: prostatic intraepithelial neoplasia.

reduced lesion odor in 90%, dry lesions in 70%, and smaller lesion size and pain mitigation in 10%.

#### 6.4. Curcumin lowers serum cholesterol and lipid peroxide levels in healthy individuals

While investigating the mechanisms of curcumin's chemopreventive effects, in another study, Kuttan and coworker

[175] monitored curcumin's effect on serum cholesterol and lipid peroxide levels in 10 healthy volunteers. Daily administration of curcumin (500 mg) for 7 days led to a significant 33% decrease in serum lipid peroxides, a 29% increase in serum HDL cholesterol, and a nearly 12% decrease in total serum cholesterol. Together, these striking findings suggest a potential chemopreventive role for curcumin in arterial diseases [175]. In Concordant with these findings are results



622 of another study in which curcumin (10 mg) administered  
623 twice a day for 28 days lowered serum LDL and increased  
624 serum HDL levels in patients with atherosclerosis [176].

#### 6.5. Curcumin may prevent gallstone formation

625 Curcumin has been evaluated for its ability to induce gall  
626 bladder emptying and thus reduce gallstone formation, a  
627 potential risk factor for gall bladder cancer. Agents that can  
628 induce the gall bladder to contract and empty itself (e.g.,  
629 erythromycin, fatty meals, and amino acids) have been  
630 shown to reduce gallstone formation. In a randomized,  
631 double-blind, crossover study involving 12 healthy volun-  
632 teers [178], 20 mg curcumin produced a positive cholekinetic  
633 effect that led to 29% contraction of the gall bladder. A  
634 subsequent study indicated that doses of 40 and 80 mg  
635 curcumin produced 50% and 72% contraction of the gall  
636 bladder volume, respectively. Together, these results suggest  
637 that curcumin can effectively induce the gall bladder to  
638 empty and thereby reduce the risk of gallstone formation and  
639 ultimately gall bladder cancer.

#### 6.6. Curcumin is effective in patients with chronic anterior uveitis and idiopathic inflammatory orbital pseudotumors

641 Curcumin's anti-inflammatory effect has also been evaluated  
642 in two rare inflammatory diseases—chronic anterior uveitis  
643 (CAU) and idiopathic inflammatory orbital pseudotumors  
644 (IOTs). In a study by Lal et al. [180] involving patients with  
645 CAU, curcumin was administered orally at a dose of 375 mg  
646 three times a day for 12 weeks. Patients were segregated into  
647 two groups: 18 patients who received curcumin alone and 14  
648 patients who, in addition to CAU, had a strong reaction to a  
649 PPD tuberculosis test and so received antitubercular treat-  
650 ment in addition to curcumin. Patients in both groups began  
651 showing improving after 2 weeks of treatment, although  
652 those in the combination therapy group had a better response  
653 rate of 86%. Moreover, at 3 years of follow-up, the recurrence  
654 rate was much lower in the combination therapy group than  
655 in the group treated with curcumin only (36% versus 55%).  
656 Although approximately one in five patients in each treat-  
657 ment group lost their vision in the follow-up period because of  
658 various complications of the primary disease (e.g., vitritis,  
659 macular edema, central venous block, cataract formation,  
660 and glaucomatous optic nerve damage), none reported any  
661 side effects of the curcumin therapy. In fact, in terms of safety  
662 and efficacy, curcumin compared favorably with the only  
663 current standard treatment for CAU (i.e., corticosteroid  
664 therapy).

665 Encouraged by this clinical study, Lal et al. [181] proceeded  
666 to evaluate curcumin as treatment for IOT and found it to  
667 be both safe and effective. In that relatively small study,  
668 eight patients took curcumin orally at a dose of 375 mg three  
669 times a day for 6–22 months and were followed up every 3  
670 months for 2 years. Although only five patients completed  
671 the study, four of them recovered completely and the fifth  
672 experienced a complete resolution of tumor-related swelling  
673 despite some residual limits on range of motion. Just  
674 as encouraging was the lack of any recurrence or side  
675 effects.

#### 6.7. Curcumin beneficially affects psoriasis

676 Curcumin has also been shown to have beneficial effects on  
677 psoriasis, another proinflammatory and potentially arthritis-  
678 inducing skin disease. In one particular study, Heng et al. [182]  
679 evaluated curcumin's antipsoriatic effects indirectly by  
680 measuring its influence on phosphorylase kinase activity.  
681 (Curcumin is a potent selective inhibitor of phosphorylase  
682 kinase, increased levels of which are considered by some to be  
683 a surrogate marker of psoriatic disease.) Phosphorylase kinase  
684 activity was assayed in four groups of 10 patients each: (i)  
685 those with active untreated psoriasis; (ii) those with resolving  
686 psoriasis treated with calcipotriol, a vitamin D3 analogue and  
687 an indirect inhibitor of phosphorylase kinase; (iii) those with  
688 resolving psoriasis treated with curcumin; and (iv) normal  
689 nonpsoriatic subjects. Phosphorylase kinase activity was  
690 highest in the patients with active untreated psoriasis,  
691 lower in the calcipotriol-treated group, even lower in the  
692 curcumin-treated group, and lowest in normal subjects.  
693 Interestingly, the decreased phosphorylase kinase activity in  
694 calcipotriol- and curcumin-treated patients was associated  
695 with corresponding decreases in the expression of keratino-  
696 cyte transferrin receptor (TRR), severity of parakeratosis, and  
697 density of epidermal CD8+ T cells.

#### 6.8. Curcumin safely exerts chemopreventive effects against multiple human cancers

698 Apparently, curcumin can also safely exert chemopreventive  
699 effects on premalignant lesions. In a prospective phase I dose-  
700 escalation study, Chen et al. [195] examined the safety,  
701 efficacy, and pharmacokinetics of curcumin in 25 patients  
702 with a variety of high-risk. Precancerous lesions (i.e., recently  
703 resected urinary bladder cancer ( $n=2$ ), arsenic Bowen's  
704 disease of the skin ( $n=6$ ), uterine cervical intraepithelial  
705 neoplasm [CIN] ( $n=4$ ), oral leukoplakia ( $n=7$ ), and intestinal  
706 metaplasia of the stomach ( $n=6$ )). Curcumin was adminis-  
707 tered to the first three patients at a starting dose of 500 mg/day  
708 for 3 months and, if no grade 2 or higher toxicities were  
709 observed, was increased to 1000, 2000, 4000, 8000, and finally  
710 12,000 mg/day. Curcumin was not toxic at doses of 8000 mg/  
711 day or lower, reaching peak serum concentrations at 1–2 h  
712 ( $0.51 \pm 0.11 \mu\text{M}$  at 4000 mg,  $0.63 \pm 0.06 \mu\text{M}$  at 6000 mg, and  
713  $1.77 \pm 1.87 \mu\text{M}$  at 8000 mg) and being gradually eliminated  
714 (principally through nonurinary routes) within 12 h. Although  
715 frank malignancies occurred despite curcumin treatment in  
716 one patient each with CIN and oral leukoplakia, a remarkable  
717 number of patients (i.e., one patient with recently resected  
718 bladder cancer, two with oral leukoplakia, one with intestinal  
719 metaplasia of the stomach, one with CIN, and two with  
720 Bowen's disease) showed histologic improvement of their  
721 precancerous lesions.

#### 6.9. Curcumin modulates biomarkers of colorectal cancer

722 Curcumin can also apparently modulate biomarkers of color-  
723 ectal cancer. In a pilot dose-escalation study in 15 patients  
724 with drug-resistant advanced colorectal cancer, Sharma et al.  
725 [160] assessed the pharmacodynamics and pharmacokinetics  
726 of a novel encapsulated turmeric extract administered at

doses ranging from 440 to 2200 mg/day for up to 4 months. (Depending on the dose, each capsule contained 36–180 mg of curcumin.) The compound's effects were measured in terms of its effects on two surrogate biomarkers (i.e., glutathione-S-transferase [GST] activity and DNA adducts formed between M(1)G and malondialdehyde) in blood cells. The compound was deemed safe and effective after the investigators observed no dose-limiting toxicity and a significant (59%) decrease in GST activity at the lowest dose (440 mg) but none at higher doses and clinically effective, and radiologically stable disease in 33% (5/15) of patients after 2–4 months of treatment.

In a subsequent dose-escalation study in a similar population, Sharma et al. [183] further explored the pharmacology of curcumin administered in capsules at daily doses ranging from 0.45 to 3.6 g daily for up to 4 months. This time, the compound's effects on leukocytes were measured in terms of three potential biomarkers: GST activity, deoxyguanosine adduct M(1)G levels, and PGE<sub>2</sub> production *ex vivo*. In a comparison of inducible PGE<sub>2</sub> production immediately before and 1 h after dosing on days 1 and 29, the highest dose (3.6 g) elicited significant decreases (62% and 57%, respectively). Consequently, the investigators chose the 3.6 g dose for further evaluation in a phase II trial in cancers outside the gastrointestinal tract.

In a subsequent and similar study, the same investigators asked whether pharmacologically active levels of curcumin could be achieved in the colorectum of colorectal cancer patients [161]. Encapsulated curcumin was administered orally at three different daily doses (3600, 1800, or 450 mg) for 7 days. Its biodistribution was then assayed by comparing curcumin levels in biopsied specimens of normal and malignant colorectal tissue obtained at diagnosis and 6–7 h after the last curcumin dose, measuring the levels of M(1)G and COX-2 protein in blood samples obtained 1 h after the last curcumin dose, and quantitating blood levels of curcumin and its metabolites by high-performance liquid chromatography and UV spectrophotometry or mass spectrometry. At the highest dose (3600 mg), the concentrations of curcumin differed between normal and malignant tissues ( $12.7 \pm 5.7$  versus  $7.7 \pm 1.8$  nmol/g). However, both normal and malignant tissues from patients so treated contained curcumin sulfate and curcumin glucuronide, and their peripheral circulation contained trace amounts of curcumin. Furthermore, the DNA adduct M(1)G was 2.5 times more abundant in cancerous tissues than in normal tissues. At the highest dose (3600 mg), curcumin lowered M<sub>1</sub>G levels (from  $4.8 \pm 2.9$  to  $2.0 \pm 1.8$  adducts per  $10^7$  nucleotides) but not COX-2 protein levels in cancerous tissues. Together, these results suggested that curcumin orally administered at a dose of 3600 mg could reach pharmacologically efficacious levels in the colorectum while at the same time being negligibly distributed outside the gut [161].

### 6.10. Curcumin helps reduce symptoms of irritable bowel syndrome

There is evidence that curcumin may help relieve symptoms of the extremely common gastric disorder known as irritable bowel syndrome (IBS). This chronic condition is characterized by abdominal pain, alterations in bowel habits and stool

frequency, and poor quality of life and appears to be causally associated with antibiotic use and inflammatory infection. In a partially blinded, randomized, pilot study in which 207 healthy adults were randomly assigned to receive either one or two tablets of a standardized turmeric extract daily for 8 weeks, IBS symptoms improved significantly after treatment [184].

In a study by another group of investigators, oral curcumin was administered in daily doses ranging from 450 to 3600 mg to 12 patients about to undergo surgery for hepatic metastases of colorectal cancer to determine whether enough of the curcumin would reach normal and malignant human liver tissue in concentrations sufficient to elicit pharmacologic activity [156]. The compound's resulting poor bioavailability (as indicated by low nanomolar levels of the parent compound and its glucuronide and sulfate conjugates in the peripheral or portal circulation) led the investigators to conclude that achieving pharmacologically effective concentrations of curcumin in the liver is not feasible.

### 6.11. Curcumin improves early renal graft function

Curcumin has also been shown to beneficially influence early kidney graft function, presumably due to its known ability to induce the activity of the antioxidant hemoxygenase-1. In a randomized, placebo-controlled trial, a combination of curcumin 480 mg and quercetin 20 mg was administered orally in capsule form to cadaveric kidney transplant recipients for 1 month, starting immediately after transplantation. The trial's 43 subjects were randomly assigned to placebo (control), low-dose (one capsule + one placebo), or high dose (two capsule) regimens [185]. Graft function was assessed in terms of delayed graft function (i.e., the need for dialysis in the first week after transplantation) and slowed graft function (i.e., serum creatinine  $>2.5$  mg/dL by post-transplantation day 10). The investigators consequently observed much better early graft function in treated patients than in controls (71% [low-dose] versus 93% [high-dose] versus 43% [controls]), no delayed graft function in any treated patients but delayed function in 14% (2/14) of controls, and significantly lower serum creatinine levels in treated patients after 2 and 30 days of treatment. They also noted significantly higher levels of urinary HO-1 in the two active treatment groups. Interestingly, however, when compared with both the low-dose and control regimens, only the high-dose regimen appeared to lower the incidence of acute graft rejection at 6 months posttransplantation (0% versus 14.3%) and reduce the incidence of tremors (13% versus 46%).

### 6.12. Curcumin improves clinical outcome in patients with tropical pancreatitis

Curcumin appears to improve the clinical outcomes of patients suffering from chronic pancreatitis, an intensely painful inflammatory condition induced by oxidative stress, by reversing lipid peroxidation. As shown in a randomized, placebo-controlled pilot study involving 20 patients with tropical pancreatitis, an oral combination of curcumin 500 mg and piperine 5 mg provided effective pain relief and beneficially modulated a pair of markers of oxidative stress

- 847 (i.e., significantly reduced malonyldialdehyde levels and  
848 increased glutathione levels in erythrocytes) [186].
- 849 **6.13. Curcumin is therapeutic in patients with**  
850 **inflammatory bowel disease**
- 851 Curcumin also appears to have beneficial therapeutic effects  
852 on inflammatory bowel disease. Marked by chronic inflam-  
853 mation of the colon and encompassing both ulcerative colitis  
854 and Crohn's disease, inflammatory bowel disease is a  
855 frequent complication of and risk factor for colorectal cancer  
856 in humans. In a preliminary open-label study based on its  
857 preclinically established anti-inflammatory and antioxidant  
858 properties, curcumin was administered to a small popula-  
859 tion of patients with previously treated ulcerative proctitis  
860 ( $n = 5$ ) or Crohn's disease ( $n = 5$ ) [187]. The five patients with  
861 ulcerative proctitis, who had been previously treated with 5-  
862 aminosalicylic acid (5ASA) compounds and (in four cases)  
863 corticosteroids, received curcumin orally at a dose of 550 mg  
864 twice daily for 1 month and then three times daily for  
865 another month. The five patients with Crohn's disease  
866 received curcumin orally at a dose of 360 mg (one capsule)  
867 three times daily for 1 month and then 360 mg (four  
868 capsules) four times daily for another 2 months. By study's  
869 end, all five cases of ulcerative proctitis had significantly  
870 improved to the point that two patients stopped taking  
871 5ASAs and two others (including one who stopped taking  
872 prednisone) reduced their 5ASA dosages. This improvement  
873 was documented in terms of a return to normal limits of the  
874 inflammatory indices of sedimentation rate and C-reactive  
875 protein (CRP) level. Meanwhile, although only four of five  
876 Crohn's disease patients completed the study, those four  
877 experienced also marked clinical improvement after curcu-  
878 min treatment, as evidenced by reductions in several indices  
879 including Crohn's disease activity index (CDAI) scores,  
880 sedimentation rate (i.e., a mean reduction of 10 mm/h,  
881 and CRP (i.e., a mean reduction of 0.1 mg/dL). Moreover,  
882 these four patients continued to show significant sympto-  
883 matic improvement (i.e., more formed stools, less frequent  
884 bowel movements, and less abdominal pain and cramping)  
885 at monthly follow-up visits. In light of these extremely  
886 encouraging findings, the investigators concluded that  
887 double-blind placebo-controlled follow-up studies were  
888 warranted.
- 889 In a subsequent randomized, double-blind, placebo-con-  
890 trolled multicenter trial [188], Hanai et al. demonstrated  
891 curcumin's ability to safely and effectively prevent the relapse  
892 of quiescent ulcerative colitis when delivered as maintenance  
893 therapy. The 89 patients enrolled in the trial were randomly  
894 assigned to a 6-month regimen of either placebo ( $n = 44$ ) or  
895 curcumin 1000 mg after breakfast and 1000 mg after dinner  
896 ( $n = 45$ ) in combination with sulfasalazine or mesalamine.  
897 After 6 months of treatment, the relapse rate among evaluable  
898 patients ( $n = 82$ ) was significantly higher in the placebo group  
899 (20.5% [8/39]), than in the curcumin-treated group (4.7% [2/43]).  
900 Curcumin also appeared to suppress disease-associated  
901 morbidity, as assessed in terms of clinical activity index  
902 (CAI) and endoscopic index (EI) scores. After an additional 6-  
903 month follow-up period, during which patients in both groups  
904 took sulfasalazine or mesalamine, another 8 curcumin-
- 905 treated patients and another 6 placebo-treated patients  
906 experienced a disease relapse.
- 907 **6.14. Curcumin reduces polyp numbers in patients with**  
908 **familial adenomatous polyposis**
- 909 Curcumin also appears to safely exert beneficial effects in  
910 patients with FAP, an autosomal-dominant disorder char-  
911 acterized by the formation of hundreds of colorectal adeno-  
912 mas and eventually the development of colorectal cancer.  
913 Typically, the growth of the adenomatous polyps is controlled  
914 in part by treatment with nonsteroidal anti-inflammatory  
915 drugs and COX-2 inhibitors, despite the considerable side  
916 effects. Therefore, in a very small clinical trial, Cruz-Correa  
917 et al. [189] evaluated curcumin's ability to induce adenoma  
918 regression in previously colectomized patients with FAP. In all  
919 five cases, combination treatment with curcumin 480 mg and  
920 quercetin 20 mg orally three times a day for a mean duration of  
921 6 months significantly decreased mean polyp number and size  
922 by 60.4% and 50.9%, respectively, without producing any  
923 noticeable toxic side effects.
- 924 **6.15. Curcumin may improve cognitive function in the**  
925 **elderly**
- 926 Despite preclinical evidence of curcumin's ability to bind  $\beta$ -  
927 amyloids and thereby reduce plaque burdens [51], there has  
928 been little, if any, supporting epidemiologic evidence of this.  
929 However, in a recent large, population-based study of 1010  
930 elderly nondemented Asians, those who consumed curry  
931 "occasionally" and "often or very often" scored significantly  
932 better on the Mini-Mental State Examination (MMSE), a  
933 established measure of cognitive function, than did those  
934 who "never or rarely" consumed curry [190]. At the least, this  
935 finding warrants further investigation of curcumin's cognitive  
936 effects.
- 937 **6.16. Curcumin may beneficially influence several cancer**  
938 **precursor conditions**
- 939 In addition to the published studies reviewed above, several  
940 other trials have been investigating curcumin's therapeutic  
941 and chemopreventive potential in certain cancer precursor  
942 conditions. One of them, a small 18-month study involving 24  
943 human subjects and still in progress, is investigating curcu-  
944 min's effect on prostatic intraepithelial neoplasia (PIN), a  
945 precursor of prostate cancer, when given in combination with  
946 a herbal product called zyflamend [191]. Another study,  
947 recently reported, found curcumin to exert beneficial effects  
948 in patients with *H. pylori* infection, a precursor of gastric cancer  
949 [192].
- 950 **6.17. Curcumin has potential in advanced pancreatic**  
951 **cancer**
- 952 Curcumin has also been examined as a single-agent in  
953 patients with advanced pancreatic cancer [196]. A dose of  
954 8 g curcumin per day was administered for 2 months. The  
955 results of this study showed that curcumin is well tolerated  
956 and a sign of biological activity found in most patients.



**Table 7 – A list of ongoing clinical trials with curcumin in patients with different diseases**

| Disease                             | Study type/design                           | Patients # | Start date     | Trial site   |
|-------------------------------------|---|------------|----------------|--|
| Colon cancer                        | Phase-I, randomized                         | 24         | Completed      | University of Michigan, Ann Arbor, USA                         |
| Colorectal cancer, ACF <sup>1</sup> | Phase-I, randomized <sup>2</sup>            | –          | Suspended      | Rockefeller University Hospital, New York, USA                 |
| Colon cancer                        | Phase-III, randomized                       | 100        | March 2006     | Tel-Aviv Sourasky Medical Center, Tel-Aviv, Israel             |
| Colorectal cancer, ACF <sup>1</sup> | Phase-II, non-randomized                    | 48         | September 2006 | University of Illinois, Chicago, USA                           |
| FAP                                 | Phase-II, randomized <sup>4</sup>           | 68         | July 2005      | University of Pennsylvania, Philadelphia, USA                  |
| FAP                                 | Phase-II, non-randomized                    | –          | November 2005  | Johns Hopkins University, Baltimore, USA                       |
| Aberrant crypt foci                 | Prevention, randomized <sup>5</sup>         | 60         | April 2004     | Cancer Institute of New Jersey, New Brunswick, USA             |
| Pancreatic cancer                   | Phase-II, non-randomized <sup>6</sup>       | 45         | July 2004      | Rambam Medical Center, Haifa, Israel                           |
| Pancreatic cancer                   | Phase-II, non-randomized                    | 50         | November 2004  | M.D. Anderson Cancer Center, Houston, USA                      |
| Pharmacokinetics                    | Treatment, non-randomized                   | 6          | August 2005    | Massachusetts General Hospital, Boston, USA                    |
| Myelodysplastic syndrome            | Phase II                                    | 30         |                | University Massachusetts, Worcester, USA (Raza A.)             |
| Alzheimer's disease                 | Phase-II, randomized                        | 33         | July 2003      | University of California Los Angeles, Los Angeles, USA         |
| Alzheimer's disease                 | Phase-I and II, randomized <sup>7</sup>     | 30         | Completed      | Chinese University of Hong Kong, Shatin, Hong Kong             |
| Multiple myeloma                    | Randomized <sup>8</sup>                     | 30         | November 2004  | M.D. Anderson Cancer Center, Houston, USA                      |
| Myelodysplastic syndrome            | Phase-I and II, non-randomized <sup>9</sup> | 50         | December 2006  | Hadassah Medical Organization, Jerusalem, Israel               |
| Psoriasis                           | Phase-II, non-randomized <sup>10</sup>      | –          | October 2005   | University of Pennsylvania, Philadelphia, USA                  |
| Epilepsy                            | Phase 1                                     | ?          | ?              | AIIMS, Delhi, India (Gupta Y.K.)                               |
| Advanced HNSCC                      | Phase II (1–8 g/day; 56 days)               | 40         | ?              | Himalyan Institute of Medical Sciences, India (Saini S.)       |
| HNSCC                               | Phase II/III DBRPC (3.6 g/day, bid)         | 300        | ?              | AIIMS, Delhi, India (Bahadur S./Ranju R./Rath G.K./Julka P.K.) |
| Cervical cancer (Stage IIb, IIIb)   | Phase II/III DBRPC (2 g/day, bid, 1 year)   | 100        | ?              | AIIMS, Delhi, India (Singh N./Jain S.K./Rath G.K./Julka P.K.)  |
| Oral premalignant lesions           | Phase II/III DBRPC (4 g/day, bid × 28 days) | 90         | ?              | Tata Memorial Cancer Center, India (D'Cruz A.)                 |
| Oral premalignant lesions           | Phase II/III DBRPC (3.6 g/day, bid)         | 96         | November 2006  | Amrita Institute, Kochi, India (Kuriakose M.A.)                |
| Oral leukoplakia                    | Phase II (curcumin gel, 3×/day, 6 month)    | 100        | ?              | Regional Cancer Center, India (Ramadas K., Pillai M.R.)        |
| Gall bladder cancer                 | Phase II (2–8 g/day)                        | 60         | ?              | BHU, India (Shukla V.K.)                                       |
| Pancreatic cancer                   | Phase II (8 g/day)                          | 40         | August 2007    | Kyoto University, Japan (Kanai M., Guha S.)                    |
| PSC                                 | Phase I (8 g/day)                           | 20         | August 2007    | Amsterdam Medical Center (Krishnadath K., Guha S.)             |
| Ulcerative colitis                  | Phase I (8 g/day)                           | 20         | August 2007    | Amsterdam Medical Center (Krishnadath K., Guha S.)             |
| Barretts Metaplasia                 | Phase I (8 g/day)                           | 20         | August 2007    | Amsterdam Medical Center (Krishnadath K., Guha S.)             |
| MGUS                                | Phase 1 (3.4 g/day)                         |            |                | St. George Hospital, Sydney (Terrance Diamond)                 |

ACF, aberrant crypt foci; DBRPC, double-blind randomized placebo-controlled; clinical trials were performed with curcumin in combination with 2. quercetin<sup>2</sup>, sulindac; 2, celecoxib; 3, 4, curcuminoids; 5, NSAIDs; 6, gemcitabine; 6, ginkgo extract; 7, bioperine; 8, coenzyme Q10; 10, curcuminoids G3 complex; 11, gemcitabine + S-1; PSC: Primary Sclerosing Cholangitis. Website: [www.clinicaltrials.gov](http://www.clinicaltrials.gov).

## 7. Ongoing clinical trials of curcumin

Enthusiasm for further studies of curcumin's chemopreventive and therapeutic effects continues to grow. Three

trials of curcumin have recently concluded, although their results have yet to be published. At least 12 active clinical trials of curcumin are ongoing in the United States, Israel, and Hong Kong (Table 7). Curcumin is being used alone in

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most of these trials and in combination with quercetin or sulindac in one. Meanwhile, chemoprevention trials of curcumin in hepatocellular carcinoma, gastric cancer, and colon cancer are ongoing in Japan. Here in the United States, several randomized and nonrandomized phase I/II trials ([www.ClinicalTrials.gov](http://www.ClinicalTrials.gov)) are investigating curcumin's effects on a range of human malignancies (e.g., colorectal cancer, aberrant crypt foci, FAP, pancreatic cancer, multiple myeloma, Alzheimer's disease, myelodysplastic syndrome, and psoriasis) when given alone or in conjunction with other natural substances or nonsteroidal anti-inflammatory drugs (NSAIDs).

Five ongoing phase I/II trials are studying curcumin's preventive and therapeutic effects on colorectal cancers in patients with FAP and ACF. Two-phase II trials are interrogating the effects of curcumin in advanced pancreatic cancers. An Israeli trial is investigating the combined effects of curcumin and gemcitabine in patients with chemotherapy-naïve, locally advanced or metastatic adenocarcinomas of the pancreas, while an exploratory clinical trial in the United States is testing the efficacy of curcumin alone in patients with unresectable or metastatic pancreatic cancers.

Two double-blind, placebo-controlled phase II trials are evaluating the efficacy, safety, and tolerability of two doses of curcumin C3 complex versus placebo in patients with mild to moderate Alzheimer's disease. An Israeli clinical trial is investigating the clinical efficacy of curcumin alone or in combination with coenzyme Q10 in patients with myelodysplastic syndrome (MDS). At M.D. Anderson Cancer Center, a pilot trial of curcumin alone or in combination with bioprine (a black pepper extract) is underway in patients with asymptomatic multiple myeloma.

## 8. Adverse effects of curcumin

Though curcumin is demonstrably bioactive and nontoxic, there are rare anecdotal reports of its deleterious side effects under certain conditions. Frank et al. [197] reported that copper-bound curcumin loses its ability to inhibit liver and kidney tumors in Cinnamon rats. Others have noted that curcumin can exhibit some blood-thinning properties such as suppression of platelet aggregation, although it remains to be established whether curcumin interacts in any way with blood-thinning drugs. Although several published studies suggest that curcumin may beneficially induce apoptosis in part through its induction of p53 expression [198], at least two other studies suggest that curcumin may instead have a deleterious, antiapoptotic effect by downregulating p53 [199,200]. Similarly, although dozens of studies indicate that curcumin potentiates the effect of chemotherapeutic agents, at least one study done in mice suggests that a curcumin-supplemented diet may inhibit the antiproliferative effects of cyclophosphamide on breast cancer growth (the investigators in that study, however, monitored tumor growth for only 93 days) [201]. There have also been reports of curcumin-induced allergic contact dermatitis [202,203] and urticaria in humans.

## 9. Conclusions

Extensive research over the last half century has made clear that most chronic illnesses can only be cured by multi-targeted, as opposed to mono-targeted, therapy [204–206] and that promiscuous targeting of a disease cell's multiple bypass mechanisms is a therapeutic virtue [207]. Consequently, agents that can modulate multiple cellular targets are now attractive objects of research. As this review has shown, curcumin is one such agent and has the potential to treat a variety of diseases. More extensive, well-controlled clinical trials are now needed to fully evaluate its potential in terms of optimal dose, route of administration, and disease targets and potential interactions with other drugs. In light of the long and established experience with curcumin as a foodstuff and as a natural medicine in humans, its low cost, its proven chemopreventive and therapeutic potential, and its pharmacological safety, curcumin is moving rapidly from the kitchen shelf toward the clinic.

## Uncited reference

[63].

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## REFERENCES

- [1] Newman DJ, Cragg GM, Snader KM. Natural products as sources of new drugs over the period 1981–2002. *J Nat Prod* 2003;66:1022–37.
- [2] Butler MS. The role of natural product chemistry in drug discovery. *J Nat Prod* 2004;67:2141–53.
- [3] Balunas MJ, Kinghorn AD. Drug discovery from medicinal plants. *Life Sci* 2005;78:431–41.
- [4] Gurib-Fakim A. Medicinal plants: traditions of yesterday and drugs of tomorrow. *Mol Aspects Med* 2006;27:1–93.
- [5] Newman DJ, Cragg GM. Natural products as sources of new drugs over the last 25 years. *J Nat Prod* 2007;70:461–77.
- [6] Mukherjee PK, Wahile A. Integrated approaches towards drug development from Ayurveda and other Indian system of medicines. *J Ethnopharmacol* 2006;103:25–35.
- [7] Kiuchi F, Goto Y, Sugimoto N, Akao N, Kondo K, Tsuda Y. Nematocidal activity of turmeric: synergistic action of curcuminoids. *Chem Pharm Bull (Tokyo)* 1993;41:1640–3.
- [8] Ravindran PN. Turmeric—the golden spice of life. *Turmeric: The genus Curcuma*. Taylor and Francis Group; 2006. p. 1–14.
- [9] Vogel and Pelletier. *J Pharm* 1818; 2:50.
- [10] Daybe FV. *Uber Curcumin*. *den Farbstoff der Curcumawurzel Ber* 1870;3:609.

- 1073 [11] Lampe V. Milobedzka. J Ver Dtsch Chem Ges 1913;46:2235. 1142
- 1074 [12] Aggarwal BB, Kumar A, Bharti AC. Anticancer potential of 1143
- 1075 curcumin: preclinical and clinical studies. Anticancer Res 1144
- 1076 2003;23:363-98. 1145
- 1077 [13] Shen L, Ji HF. Theoretical study on physicochemical 1146
- 1078 properties of curcumin. Spectrochim Acta A Mol Biomol 1147
- 1079 Spectrosc 2007;67:619-23. 1148
- 1080 [14] Bernabe-Pineda M, Ramirez-Silva MT, Romero-Romo M, 1149
- 1081 Gonzalez-Vergara E, Rojas-Hernandez A. Determination of 1150
- 1082 acidity constants of curcumin in aqueous solution and 1151
- 1083 apparent rate constant of its decomposition. Spectrochim 1152
- 1084 Acta A Mol Biomol Spectrosc 2004;60:1091-7. 1153
- 1085 [15] Wang YJ, Pan MH, Cheng AL, Lin LI, Ho YS, Hsieh CY, et al. 1154
- 1086 Stability of curcumin in buffer solutions and 1155
- 1087 characterization of its degradation products. J Pharm 1156
- 1088 Biomed Anal 1997;15:1867-76. 1157
- 1089 [16] Tonnesen HH, Karlsen J. Studies on curcumin and 1158
- 1090 curcuminoids. VI. Kinetics of curcumin degradation in 1159
- 1091 aqueous solution. Z Lebensm Unters Forsch 1985;180: 1160
- 1092 402-4. 1161
- 1093 [17] Oetari S, Sudibyo M, Commandeur JN, Samhoedi R, 1162
- 1094 Vermeulen NP. Effects of curcumin on cytochrome P450 1163
- 1095 and glutathione-S-transferase activities in rat liver. 1164
- 1096 Biochem Pharmacol 1996;51:39-45. 1165
- 1097 [18] Pan MH, Huang TM, Lin JK. Biotransformation of curcumin 1166
- 1098 through reduction and glucuronidation in mice. Drug 1167
- 1099 Metab Dispos 1999;27:486-94. 1168
- 1100 [19] Somporn P, Phisalaphong C, Nakornchai S, Unchern S, 1169
- 1101 Morales NP. Comparative antioxidant activities of 1170
- 1102 curcumin and its demethoxy and hydrogenated 1171
- 1103 derivatives. Biol Pharm Bull 2007;30:74-8. 1172
- 1104 [20] Pari L, Murugan P. Tetrahydrocurcumin prevents brain 1173
- 1105 lipid peroxidation in streptozotocin-induced diabetic rats. 1174
- 1106 J Med Food 2007;10:323-9. 1175
- 1107 [21] Murugan P, Pari L. Antioxidant effect of 1176
- 1108 tetrahydrocurcumin in streptozotocin-nicotinamide 1177
- 1109 induced diabetic rats. Life Sci 2006;79:1720-8. 1178
- 1110 [22] Tomren MA, Masson M, Loftsson T, Tonnesen HH. Studies 1179
- 1111 on curcumin and curcuminoids XXXI. Symmetric and 1180
- 1112 asymmetric curcuminoids: stability, activity and 1181
- 1113 complexation with cyclodextrin. Int J Pharm 2007;338:27- 1182
- 1114 34. 1183
- 1115 [23] Govindarajan VS. Turmeric-chemistry, technology, and 1184
- 1116 quality. Crit Rev Food Sci Nutr 1980;12:199-301. 1185
- 1117 [24] Ammon HP, Wahl MA. Pharmacology of *Curcuma longa*. 1186
- 1118 Planta Med 1991;57:1-7. 1187
- 1119 [25] Araujo CC, Leon LL. Biological activities of *Curcuma longa* 1188
- 1120 L.. Mem Inst Oswaldo Cruz 2001;96:723-8. 1189
- 1121 [26] Aggarwal BB, Takada Y, Oommen OV. From 1190
- 1122 chemoprevention to chemotherapy: common targets and 1191
- 1123 common goals. Expert Opin Investig Drugs 2004;13: 1192
- 1124 1327-38. 1193
- 1125 [27] Sreejayan, Rao MN. Nitric oxide scavenging by 1194
- 1126 curcuminoids. J Pharm Pharmacol 1997;49:105-7. 1195
- 1127 [28] Brouet I, Ohshima H. Curcumin, an anti-tumour promoter 1196
- 1128 and anti-inflammatory agent, inhibits induction of nitric 1197
- 1129 oxide synthase in activated macrophages. Biochem 1198
- 1130 Biophys Res Commun 1995;206:533-40. 1199
- 1131 [29] Dikshit M, Rastogi L, Shukla R, Srimal RC. Prevention of 1200
- 1132 ischaemia-induced biochemical changes by curcumin & 1201
- 1133 quinidine in the cat heart. Indian J Med Res 1995;101: 1202
- 1134 31-5. 1203
- 1135 [30] Rao CV, Rivenson A, Simi B, Reddy BS. Chemoprevention 1204
- 1136 of colon carcinogenesis by dietary curcumin, a naturally 1205
- 1137 occurring plant phenolic compound. Cancer Res 1206
- 1138 1995;55:259-66. 1207
- 1139 [31] Limtrakul P, Lipigorngoson S, Namwong O, Apisariyakul A, 1208
- 1140 Dunn FW. Inhibitory effect of dietary curcumin on skin 1209
- 1141 carcinogenesis in mice. Cancer Lett 1997;116:197-203. 1210
- [32] Kiso Y, Suzuki Y, Watanabe N, Oshima Y, Hikino H. 1142
- Antihepatotoxic principles of *Curcuma longa* rhizomes. 1143
- Planta Med 1983;49:185-7. 1144
- [33] Srivastava R, Dikshit M, Srimal RC, Dhawan BN. Anti- 1145
- thrombotic effect of curcumin. Thromb Res 1985;40:413-7. 1146
- [34] Nirmala C, Puvanakrishnan R. Protective role of curcumin 1147
- against isoproterenol induced myocardial infarction in 1148
- rats. Mol Cell Biochem 1996;159:85-93. 1149
- [35] Venkatesan N. Curcumin attenuation of acute adriamycin 1150
- myocardial toxicity in rats. Br J Pharmacol 1998;124:425-7. 1151
- [36] Srinivasan M. Effect of curcumin on blood sugar as seen in 1152
- a diabetic subject. Indian J Med Sci 1972;26:269-70. 1153
- [37] Babu PS, Srinivasan K. Influence of dietary curcumin and 1154
- cholesterol on the progression of experimentally induced 1155
- diabetes in albino rat. Mol Cell Biochem 1995;152:13-21. 1156
- [38] Arun N, Nalini N. Efficacy of turmeric on blood sugar and 1157
- polyol pathway in diabetic albino rats. Plant Foods Hum 1158
- Nutr 2002;57:41-52. 1159
- [39] Deodhar SD, Sethi R, Srimal RC. Preliminary study on 1160
- antirheumatic activity of curcumin (diferuloyl methane). 1161
- Indian J Med Res 1980;71:632-4. 1162
- [40] Shankar TN, Shantha NV, Ramesh HP, Murthy IA, Murthy 1163
- VS. Toxicity studies on turmeric (*Curcuma longa*): acute 1164
- toxicity studies in rats, guineapigs & monkeys. Indian J 1165
- Exp Biol 1980;18:73-5. 1166
- [41] Qureshi S, Shah AH, Ageel AM. Toxicity studies on *Alpinia* 1167
- galanga* and *Curcuma longa*. Planta Med 1992;58:124-7. 1168
- [42] Lao CD, Ruffin MT, Normolle D, Heath DD, Murray SI, 1169
- Bailey JM, et al. Dose escalation of a curcuminoid 1170
- formulation. BMC Complement Altern Med 2006;6:10. 1171
- [43] Aggarwal BB, Bhatt ID, Ichikawa H, Ahn KS, Sethi G, 1172
- Sandur SK, et al. Curcumin-biological and medicinal 1173
- properties. Turmeric: the genus *Curcuma*. Taylor and 1174
- Francis Group; 2006. p. 297-368. 1175
- [44] Shishodia S, Singh T, Chaturvedi MM. Modulation of 1176
- transcription factors by curcumin. Adv Exp Med Biol 1177
- 2007;595:127-48. 1178
- [45] Pulla Reddy AC, Sudharshan E, Appu Rao AG, Lokesh BR. 1179
- Interaction of curcumin with human serum albumin-a 1180
- spectroscopic study. Lipids 1999;34:1025-9. 1181
- [46] Zsila F, Bikadi Z, Simonyi M. Unique, pH-dependent 1182
- biphasic band shape of the visible circular dichroism of 1183
- curcumin-serum albumin complex. Biochem Biophys Res 1184
- Commun 2003;301:776-82. 1185
- [47] Barik A, Priyadarsini KI, Mohan H. Photophysical studies 1186
- on binding of curcumin to bovine serum albumins. 1187
- Photochem Photobiol 2003;77:597-603. 1188
- [48] Wang F, Yang J, Wu X, Liu S. Study of the interaction of 1189
- proteins with curcumin and SDS and its analytical 1190
- application. Spectrochim Acta A Mol Biomol Spectrosc 1191
- 2005;61:2650-6. 1192
- [49] Kunwar A, Barik A, Pandey R, Priyadarsini KI. Transport of 1193
- liposomal and albumin loaded curcumin to living cells: an 1194
- absorption and fluorescence spectroscopic study. Biochim 1195
- Biophys Acta 2006;1760:1513-20. 1196
- [50] Zsila F, Bikadi Z, Simonyi M. Induced circular dichroism 1197
- spectra reveal binding of the antiinflammatory curcumin 1198
- to human alpha1-acid glycoprotein. Bioorg Med Chem 1199
- 2004;12:3239-45. 1200
- [51] Yang F, Lim GP, Begum AN, Ubeda OJ, Simmons MR, 1201
- Ambegaokar SS, et al. Curcumin inhibits formation of 1202
- amyloid beta oligomers and fibrils, binds plaques, and 1203
- reduces amyloid in vivo. J Biol Chem 2005;280:5892-901. 1204
- [52] Logan-Smith MJ, Lockyer PJ, East JM, Lee AG. Curcumin, a 1205
- molecule that inhibits the Ca<sup>2+</sup>-ATPase of sarcoplasmic 1206
- reticulum but increases the rate of accumulation of Ca<sup>2+</sup>. J 1207
- Biol Chem 2001;276:46905-11. 1208
- [53] Bilmen JG, Khan SZ, Javed MH, Michelangeli F. Inhibition 1209
- of the SERCA Ca<sup>2+</sup> pumps by curcumin. Curcumin 1210

- 1211 putatively stabilizes the interaction between the  
1212 nucleotide-binding and phosphorylation domains in the  
1213 absence of ATP. *Eur J Biochem* 2001;268:6318–27.
- 1214 [54] Reddy S, Aggarwal BB. Curcumin is a non-competitive and  
1215 selective inhibitor of phosphorylase kinase. *FEBS Lett*  
1216 1994;341:19–22.
- 1217 [55] Shim JS, Kim JH, Cho HY, Yum YN, Kim SH, Park HJ, et al.  
1218 Irreversible inhibition of CD13/aminopeptidase N by the  
1219 antiangiogenic agent curcumin. *Chem Biol* 2003;10:  
1220 695–704.
- 1221 [56] Takeuchi T, Ishidoh T, Iijima H, Kuriyama I, Shimazaki N,  
1222 Koiwai O, et al. Structural relationship of curcumin  
1223 derivatives binding to the BRCT domain of human DNA  
1224 polymerase lambda. *Genes Cells* 2006;11:223–35.
- 1225 [57] Leu TH, Su SL, Chuang YC, Maa MC. Direct inhibitory  
1226 effect of curcumin on Src and focal adhesion kinase  
1227 activity. *Biochem Pharmacol* 2003;66:2323–31.
- 1228 [58] Awasthi S, Pandya U, Singhal SS, Lin JT, Thiviyanathan  
1229 V, Seifert Jr WE, et al. Curcumin–glutathione  
1230 interactions and the role of human glutathione-S-  
1231 transferase P1-1. *Chem Biol Interact* 2000;128:19–38.
- 1232 [59] Wortelboer HM, Usta M, van der Velde AE, Boersma MG,  
1233 Spenkeliink B, van Zanden JJ, et al. Interplay between MRP  
1234 inhibition and metabolism of MRP inhibitors: the case of  
1235 curcumin. *Chem Res Toxicol* 2003;16:1642–51.
- 1236 [60] Iersel ML, Ploemen JP, Struik I, van Amersfoort C, Keyzer  
1237 AE, Schefferlie JG, et al. Inhibition of glutathione-S-  
1238 transferase activity in human melanoma cells by  
1239 alpha,beta-unsaturated carbonyl derivatives. Effects of  
1240 acrolein, cinnamaldehyde, citral, crotonaldehyde,  
1241 curcumin, ethacrynic acid, and *trans*-2-hexenal. *Chem Biol*  
1242 *Interact* 1996;102:117–32.
- 1243 [61] Jung Y, Xu W, Kim H, Ha N, Neckers L. Curcumin-induced  
1244 degradation of ErbB2: a role for the E3 ubiquitin ligase  
1245 CHIP and the Michael reaction acceptor activity of  
1246 curcumin. *Biochim Biophys Acta* 2007;1773:383–90.
- 1247 [62] Baum L, Ng A. Curcumin interaction with copper and iron  
1248 suggests one possible mechanism of action in Alzheimer's  
1249 disease animal models. *J Alzheimers Dis* 2004;6:367–77.  
1250 discussion 443–9.
- 1251 [63] Ishihara M, Sakagami H. Re-evaluation of cytotoxicity and  
1252 iron chelation activity of three beta-diketones by  
1253 semiempirical molecular orbital method. *In Vivo*  
1254 2005;19:119–23.
- 1255 [64] Jankun J, Aleem AM, Malgorzewicz S, Szkudlarek M,  
1256 Zawadzki MI, Dewitt DL, et al. Synthetic curcuminoids  
1257 modulate the arachidonic acid metabolism of human  
1258 platelet 12-lipoxygenase and reduce sprout formation of  
1259 human endothelial cells. *Mol Cancer Ther* 2006;5:1371–82.
- 1260 [65] Skrzypczak-Jankun E, Zhou K, McCabe NP, Selman SH,  
1261 Jankun J. Structure of curcumin in complex with  
1262 lipoxygenase and its significance in cancer. *Int J Mol Med*  
1263 2003;12:17–24.
- 1264 [66] Gupta KK, Bharne SS, Rathinasamy K, Naik NR, Panda D.  
1265 Dietary antioxidant curcumin inhibits microtubule  
1266 assembly through tubulin binding. *FEBS J* 2006;273:5320–  
1267 32.
- 1268 [67] Zsila F, Bikadi Z, Simonyi M. Circular dichroism  
1269 spectroscopic studies reveal pH dependent binding of  
1270 curcumin in the minor groove of natural and synthetic  
1271 nucleic acids. *Org Biomol Chem* 2004;2:2902–10.
- 1272 [68] Romiti N, Tongiani R, Cervelli F, Chieli E. Effects of  
1273 curcumin on P-glycoprotein in primary cultures of rat  
1274 hepatocytes. *Life Sci* 1998;62:2349–58.
- 1275 [69] Anuchapreeda S, Leechanachai P, Smith MM, Ambudkar  
1276 SV, Limtrakul PN. Modulation of P-glycoprotein  
1277 expression and function by curcumin in multidrug-  
1278 resistant human KB cells. *Biochem Pharmacol*  
1279 2002;64:573–82.
- [70] Chearwae W, Anuchapreeda S, Nandigama K, Ambudkar  
1281 SV, Limtrakul P. Biochemical mechanism of modulation of  
1282 human P-glycoprotein (ABCB1) by curcumin I, II, and III  
1283 purified from Turmeric powder. *Biochem Pharmacol*  
1284 2004;68:2043–52.
- [71] Fang J, Lu J, Holmgren A. Thioredoxin reductase is  
1285 irreversibly modified by curcumin: a novel molecular  
1286 mechanism for its anticancer activity. *J Biol Chem*  
1287 2005;280:25284–90.
- [72] Martin-Cordero C, Lopez-Lazaro M, Galvez M, Ayuso MJ.  
1289 Curcumin as a DNA topoisomerase II poison. *J Enzyme*  
1290 *Inhib Med Chem* 2003;18:505–9.
- [73] Mullally JE, Fitzpatrick FA. Pharmacophore model for  
1292 novel inhibitors of ubiquitin isopeptidases that induce  
1293 p53-independent cell death. *Mol Pharmacol* 2002;62:351–8.
- [74] Began G, Sudharshan E, Appu Rao AG. Inhibition of  
1295 lipoxygenase 1 by phosphatidylcholine micelles-bound  
1296 curcumin. *Lipids* 1998;33:1223–8.
- [75] Lengyel E, Sawada K, Salgia R. Tyrosine kinase mutations  
1298 in human cancer. *Curr Mol Med* 2007;7:77–84.
- [76] Tikhomirov O, Carpenter G. Identification of ErbB-2 kinase  
1300 domain motifs required for geldanamycin-induced  
1301 degradation. *Cancer Res* 2003;63:39–43.
- [77] Aoki H, Takada Y, Kondo S, Sawaya R, Aggarwal B, Kondo  
1303 Y. Evidence that curcumin suppresses the growth of  
1304 malignant gliomas in vitro and in vivo through induction  
1305 of autophagy: role of Akt and ERK signaling pathways. *Mol*  
1306 *Pharmacol* 2007.
- [78] Grosser T. The pharmacology of selective inhibition of  
1308 COX-2. *Thromb Haemost* 2006;96:393–400.
- [79] Mitra A, Chakrabarti J, Banerji A, Chatterjee A, Das BR.  
1310 Curcumin, a potential inhibitor of MMP-2 in human  
1311 laryngeal squamous carcinoma cells HEP2. *J Environ*  
1312 *Pathol Toxicol Oncol* 2006;25:679–90.
- [80] Cho JW, Lee KS, Kim CW. Curcumin attenuates the  
1314 expression of IL-1beta, IL-6, and TNF-alpha as well as  
1315 cyclin E in TNF-alpha-treated HaCaT cells; NF-kappaB and  
1316 MAPKs as potential upstream targets. *Int J Mol Med*  
1317 2007;19:469–74.
- [81] Aggarwal S, Ichikawa H, Takada Y, Sandur SK, Shishodia  
1319 S, Aggarwal BB. Curcumin (diferuloylmethane) down-  
1320 regulates expression of cell proliferation and  
1321 antiapoptotic and metastatic gene products through  
1322 suppression of I kappa B alpha kinase and Akt activation.  
1323 *Mol Pharmacol* 2006;69:195–206.
- [82] Li M, Zhang Z, Hill DL, Wang H, Zhang R. Curcumin, a  
1325 dietary component, has anticancer, chemosensitization,  
1326 and radiosensitization effects by down-regulating the  
1327 MDM2 oncogene through the PI3K/mTOR/ETS2 pathway.  
1328 *Cancer Res* 2007;67:1988–96.
- [83] Song G, Mao YB, Cai QF, Yao LM, Ouyang GL, Bao SD.  
1330 Curcumin induces human HT-29 colon adenocarcinoma  
1331 cell apoptosis by activating p53 and regulating apoptosis-  
1332 related protein expression. *Braz J Med Biol Res*  
1333 2005;38:1791–8.
- [84] Lontas A, Yeger H. Curcumin and resveratrol induce  
1335 apoptosis and nuclear translocation and activation of  
1336 p53 in human neuroblastoma. *Anticancer Res* 2004;24:  
1337 987–98.
- [85] Gaedeke J, Noble NA, Border WA. Curcumin blocks fibrosis  
1339 in anti-Thy 1 glomerulonephritis through up-regulation of  
1340 heme oxygenase 1. *Kidney Int* 2005;68:2042–9.
- [86] McNally SJ, Harrison EM, Ross JA, Garden OJ, Wigmore SJ.  
1342 Curcumin induces heme oxygenase 1 through generation  
1343 of reactive oxygen species, p38 activation and  
1344 phosphatase inhibition. *Int J Mol Med* 2007;19:165–72.
- [87] Rao CV, Simi B, Reddy BS. Inhibition by dietary curcumin  
1346 of azoxymethane-induced ornithine decarboxylase,  
1347 tyrosine protein kinase, arachidonic acid metabolism and  
1348



- 1349 aberrant crypt foci formation in the rat colon.  
1350 Carcinogenesis 1993;14:2219-25.
- 1351 [88] Huang MT, Lou YR, Ma W, Newmark HL, Reuhl KR, Conney  
1352 AH. Inhibitory effects of dietary curcumin on forestomach,  
1353 duodenal, and colon carcinogenesis in mice. Cancer Res  
1354 1994;54:5841-7.
- 1355 [89] Kim JM, Araki S, Kim DJ, Park CB, Takasuka N, Baba-  
1356 Toriyama H, et al. Chemopreventive effects of carotenoids  
1357 and curcumins on mouse colon carcinogenesis after 1,2-  
1358 dimethylhydrazine initiation. Carcinogenesis 1998;19:  
1359 81-5.
- 1360 [90] Rao CV, Rivenson A, Simi B, Reddy BS. Chemoprevention  
1361 of colon cancer by dietary curcumin. Ann N Y Acad Sci  
1362 1995;768:201-4.
- 1363 [91] Kawamori T, Lubet R, Steele VE, Kelloff GJ, Kaskey RB, Rao  
1364 CV, et al. Chemopreventive effect of curcumin, a naturally  
1365 occurring anti-inflammatory agent, during the promotion/  
1366 progression stages of colon cancer. Cancer Res  
1367 1999;59:597-601.
- 1368 [92] Collett GP, Robson CN, Mathers JC, Campbell FC. Curcumin  
1369 modifies Apc(min) apoptosis resistance and inhibits 2-  
1370 amino 1-methyl-6-phenylimidazo[4,5-b]pyridine (PhIP)  
1371 induced tumour formation in Apc(min) mice.  
1372 Carcinogenesis 2001;22:821-5.
- 1373 [93] Pereira MA, Grubbs CJ, Barnes LH, Li H, Olson GR, Eto I,  
1374 et al. Effects of the phytochemicals, curcumin and  
1375 quercetin, upon azoxymethane-induced colon cancer and  
1376 7,12-dimethylbenz[a]anthracene-induced mammary  
1377 cancer in rats. Carcinogenesis 1996;17:1305-11.
- 1378 [94] Kwon Y, Malik M, Magnuson BA. Inhibition of colonic  
1379 aberrant crypt foci by curcumin in rats is affected by age.  
1380 Nutr Cancer 2004;48:37-43.
- 1381 [95] Shpitz B, Giladi N, Sagiv E, Lev-Ari S, Liberman E, Kazanov  
1382 D, et al. Celecoxib and curcumin additively inhibit the  
1383 growth of colorectal cancer in a rat model. Digestion  
1384 2006;74:140-4.
- 1385 [96] Sugimoto K, Hanai H, Tozawa K, Aoshi T, Uchijima M,  
1386 Nagata T, et al. Curcumin prevents and ameliorates  
1387 trinitrobenzene sulfonic acid-induced colitis in mice.  
1388 Gastroenterology 2002;123:1912-22.
- 1389 [97] Salh B, Assi K, Templeman V, Parhar K, Owen D, Gomez-  
1390 Munoz A, et al. Curcumin attenuates DNB-induced murine  
1391 colitis. Am J Physiol Gastrointest Liver Physiol  
1392 2003;285:G235-43.
- 1393 [98] Ukil A, Maity S, Karmakar S, Datta N, Vedasiromoni JR, Das  
1394 PK. Curcumin, the major component of food flavour  
1395 turmeric, reduces mucosal injury in trinitrobenzene  
1396 sulphonic acid-induced colitis. Br J Pharmacol  
1397 2003;139:209-18.
- 1398 [99] Venkataranganna MV, Rafiq M, Gopumadhavan S, Peer G,  
1399 Babu UV, Mitra SK. NCB-02 (standardized Curcumin  
1400 preparation) protects dinitrochlorobenzene-induced  
1401 colitis through down-regulation of NFkappa-B and iNOS.  
1402 World J Gastroenterol 2007;13:1103-7.
- 1403 [100] Ushida J, Sugie S, Kawabata K, Pham QV, Tanaka T, Fujii K,  
1404 et al. Chemopreventive effect of curcumin on N-  
1405 nitrosomethylbenzylamine-induced esophageal  
1406 carcinogenesis in rats. Jpn J Cancer Res 2000;91:  
1407 893-8.
- 1408 [101] Huang MT, Deschner EE, Newmark HL, Wang ZY, Ferraro  
1409 TA, Conney AH. Effect of dietary curcumin and ascorbyl  
1410 palmitate on azoxymethanol-induced colonic epithelial  
1411 cell proliferation and focal areas of dysplasia. Cancer Lett  
1412 1992;64:117-21.
- 1413 [102] Perkins S, Verschoyle RD, Hill K, Parveen I, Threadgill MD,  
1414 Sharma RA, et al. Chemopreventive efficacy and  
1415 pharmacokinetics of curcumin in the min/+ mouse, a  
1416 model of familial adenomatous polyposis. Cancer  
1417 Epidemiol Biomarkers Prev 2002;11:535-40.
- [103] Azuine MA, Bhide SV. Chemopreventive effect of turmeric  
1418 against stomach and skin tumors induced by chemical  
1419 carcinogens in Swiss mice. Nutr Cancer 1992;17:77-83.  
1420
- [104] Singh SV, Hu X, Srivastava SK, Singh M, Xia H, Orchard JL,  
1421 et al. Mechanism of inhibition of benzo[a]pyrene-induced  
1422 forestomach cancer in mice by dietary curcumin.  
1423 Carcinogenesis 1998;19:1357-60.  
1424
- [105] Nagabhushan M, Bhide SV. Curcumin as an inhibitor of  
1425 cancer. J Am Coll Nutr 1992;11:192-8.  
1426
- [106] Ikezaki S, Nishikawa A, Furukawa F, Kudo K, Nakamura H,  
1427 Tamura K, et al. Chemopreventive effects of curcumin on  
1428 glandular stomach carcinogenesis induced by N-methyl-  
1429 N'-nitro-N-nitrosoguanidine and sodium chloride in rats.  
1430 Anticancer Res 2001;21:3407-11.  
1431
- [107] Chuang SE, Cheng AL, Lin JK, Kuo ML. Inhibition by  
1432 curcumin of diethylnitrosamine-induced hepatic  
1433 hyperplasia, inflammation, cellular gene products and  
1434 cell-cycle-related proteins in rats. Food Chem Toxicol  
1435 2000;38:991-5.  
1436
- [108] Hecht SS, Kenney PM, Wang M, Trushin N, Agarwal S, Rao  
1437 AV, et al. Evaluation of butylated hydroxyanisole, myo-  
1438 inositol, curcumin, esculetin, resveratrol and lycopene as  
1439 inhibitors of benzo[a]pyrene plus 4-(methylnitrosamino)-  
1440 1-(3-pyridyl)-1-butanone-induced lung tumorigenesis in  
1441 A/J mice. Cancer Lett 1999;137:123-30.  
1442
- [109] Huang MT, Lou YR, Xie JG, Ma W, Lu YP, Yen P, et al. Effect  
1443 of dietary curcumin and dibenzoylmethane on formation  
1444 of 7,12-dimethylbenz[a]anthracene-induced mammary  
1445 tumors and lymphomas/leukemias in Sencar mice.  
1446 Carcinogenesis 1998;19:1697-700.  
1447
- [110] Singletary K, MacDonald C, Wallig M, Fisher C. Inhibition  
1448 of 7,12-dimethylbenz[a]anthracene (DMBA)-induced  
1449 mammary tumorigenesis and DMBA-DNA adduct  
1450 formation by curcumin. Cancer Lett 1996;103:137-41.  
1451
- [111] Deshpande SS, Ingle AD, Maru GB. Chemopreventive  
1452 efficacy of curcumin-free aqueous turmeric extract in  
1453 7,12-dimethylbenz[a]anthracene-induced rat mammary  
1454 tumorigenesis. Cancer Lett 1998;123:35-40.  
1455
- [112] Inano H, Onoda M, Inafuku N, Kubota M, Kamada Y,  
1456 Osawa T, et al. Chemoprevention by curcumin during the  
1457 promotion stage of tumorigenesis of mammary gland in  
1458 rats irradiated with gamma-rays. Carcinogenesis  
1459 1999;20:1011-8.  
1460
- [113] Inano H, Onoda M. Radioprotective action of curcumin  
1461 extracted from *Curcuma longa* LINN: inhibitory effect on  
1462 formation of urinary 8-hydroxy-2'-deoxyguanosine,  
1463 tumorigenesis, but not mortality, induced by gamma-ray  
1464 irradiation. Int J Radiat Oncol Biol Phys 2002;53:735-43.  
1465
- [114] Lin CC, Ho CT, Huang MT. Mechanistic studies on the  
1466 inhibitory action of dietary dibenzoylmethane, a beta-  
1467 diketone analogue of curcumin, on 7,12-  
1468 dimethylbenz[a]anthracene-induced mammary  
1469 tumorigenesis. Proc Natl Sci Counc Repub China B  
1470 2001;25:158-65.  
1471
- [115] Lin CC, Lu YP, Lou YR, Ho CT, Newmark HH, MacDonald C,  
1472 et al. Inhibition by dietary dibenzoylmethane of  
1473 mammary gland proliferation, formation of DMBA-DNA  
1474 adducts in mammary glands, and mammary  
1475 tumorigenesis in Sencar mice. Cancer Lett 2001;168:  
1476 125-32.  
1477
- [116] Azuine MA, Bhide SV. Protective single/combined  
1478 treatment with betel leaf and turmeric against methyl  
1479 (acetoxymethyl) nitrosamine-induced hamster oral  
1480 carcinogenesis. Int J Cancer 1992;51:412-5.  
1481
- [117] Tanaka T, Makita H, Ohnishi M, Hirose Y, Wang A, Mori H,  
1482 et al. Chemoprevention of 4-nitroquinoline 1-oxide-  
1483 induced oral carcinogenesis by dietary curcumin and  
1484 hesperidin: comparison with the protective effect of beta-  
1485 carotene. Cancer Res 1994;54:4653-9.  
1486



- 1487 [118] Imaida K, Tamano S, Kato K, Ikeda Y, Asamoto M, 1556  
 1488 Takahashi S, et al. Lack of chemopreventive effects of 1557  
 1489 lycopene and curcumin on experimental rat prostate 1558  
 1490 carcinogenesis. *Carcinogenesis* 2001;22:467-72. 1559  
 1491 [119] Ishizaki C, Oguro T, Yoshida T, Wen CQ, Sueki H, Iijima M. 1560  
 1492 Enhancing effect of ultraviolet A on ornithine 1561  
 1493 decarboxylase induction and dermatitis evoked by 12-*o*- 1562  
 1494 tetradecanoylphorbol-13-acetate and its inhibition by 1563  
 1495 curcumin in mouse skin. *Dermatology* 1996;193:311-7. 1564  
 1496 [120] Huang MT, Smart RC, Wong CQ, Conney AH. Inhibitory 1565  
 1497 effect of curcumin, chlorogenic acid, caffeic acid, and 1566  
 1498 ferulic acid on tumor promotion in mouse skin by 12-*o*- 1567  
 1499 tetradecanoylphorbol-13-acetate. *Cancer Res* 1568  
 1500 1988;48:5941-6. 1569  
 1501 [121] Lu YP, Chang RL, Huang MT, Conney AH. Inhibitory effect 1570  
 1502 of curcumin on 12-*o*-tetradecanoylphorbol-13-acetate- 1571  
 1503 induced increase in ornithine decarboxylase mRNA in 1572  
 1504 mouse epidermis. *Carcinogenesis* 1993;14:293-7. 1573  
 1505 [122] Huang MT, Ma W, Lu YP, Chang RL, Fisher C, Manchand 1574  
 1506 PS, et al. Effects of curcumin, demethoxycurcumin, 1575  
 1507 bisdemethoxycurcumin and tetrahydrocurcumin on 12-*o*- 1576  
 1508 tetradecanoylphorbol-13-acetate-induced tumor 1577  
 1509 promotion. *Carcinogenesis* 1995;16:2493-7. 1578  
 1510 [123] Huang MT, Ma W, Yen P, Xie JG, Han J, Frenkel K, et al. 1579  
 1511 Inhibitory effects of topical application of low doses of 1580  
 1512 curcumin on 12-*o*-tetradecanoylphorbol-13-acetate- 1581  
 1513 induced tumor promotion and oxidized DNA bases in 1582  
 1514 mouse epidermis. *Carcinogenesis* 1997;18:83-8. 1583  
 1515 [124] Soudamini KK, Kuttan R. Inhibition of chemical 1584  
 1516 carcinogenesis by curcumin. *J Ethnopharmacol* 1585  
 1517 1989;27:227-33. 1586  
 1518 [125] Takaba K, Hirose M, Yoshida Y, Kimura J, Ito N, Shirai T. 1587  
 1519 Effects of *n*-tritriacontane-16,18-dione, curcumin, 1588  
 1520 chlorophyllin, dihydroguaiaretic acid, tannic acid and 1589  
 1521 phytic acid on the initiation stage in a rat multi-organ 1590  
 1522 carcinogenesis model. *Cancer Lett* 1997;113:39-46. 1591  
 1523 [126] Huang MT, Lysz T, Ferraro T, Abidi TF, Laskin JD, Conney 1592  
 1524 AH. Inhibitory effects of curcumin on in vitro 1593  
 1525 lipoxygenase and cyclooxygenase activities in mouse 1594  
 1526 epidermis. *Cancer Res* 1991;51:813-9. 1595  
 1527 [127] Mahmoud NN, Carothers AM, Grunberger D, Bilinski RT, 1596  
 1528 Churchill MR, Martucci C, et al. Plant phenolics decrease 1597  
 1529 intestinal tumors in an animal model of familial 1598  
 1530 adenomatous polyposis. *Carcinogenesis* 2000;21:921-7. 1599  
 1531 [128] Kwon Y, Magnuson BA. Effect of azoxymethane and 1600  
 1532 curcumin on transcriptional levels of cyclooxygenase-1 1601  
 1533 and -2 during initiation of colon carcinogenesis. *Scand J* 1602  
 1534 *Gastroenterol* 2007;42:72-80. 1603  
 1535 [129] Sreepriya M, Bali G. Chemopreventive effects of embelin 1604  
 1536 and curcumin against *N*-nitrosodiethylamine/ 1605  
 1537 phenobarbital-induced hepatocarcinogenesis in Wistar 1606  
 1538 rats. *Fitoterapia* 2005;76:549-55. 1607  
 1539 [130] Sreepriya M, Bali G. Effects of administration of Embelin 1608  
 1540 and Curcumin on lipid peroxidation, hepatic glutathione 1609  
 1541 antioxidant defense and hematopoietic system during *N*- 1610  
 1542 nitrosodiethylamine/phenobarbital-induced 1611  
 1543 hepatocarcinogenesis in Wistar rats. *Mol Cell Biochem* 1612  
 1544 2006;284:49-55. 1613  
 1545 [131] Kalpana C, Rajasekharan KN, Menon VP. Modulatory 1614  
 1546 effects of curcumin and curcumin analog on circulatory 1615  
 1547 lipid profiles during nicotine-induced toxicity in Wistar 1616  
 1548 rats. *J Med Food* 2005;8:246-50. 1617  
 1549 [132] Mahady GB, Pendlan SL, Yun G, Lu ZZ. Turmeric (*Curcuma* 1618  
 1550 *longa*) and curcumin inhibit the growth of *Helicobacter* 1619  
 1551 *pylori*, a group 1 carcinogen. *Anticancer Res* 2002;22:4179- 1620  
 1552 81. 1621  
 1553 [133] Kuttan R, Bhanumathy P, Nirmala K, George MC. Potential 1622  
 1554 anticancer activity of turmeric (*Curcuma longa*). *Cancer Lett* 1623  
 1555 1985;29:197-202. 1624
- [134] Ruby AJ, Kuttan G, Babu KD, Rajasekharan KN, Kuttan R. 1556  
 Anti-tumour and antioxidant activity of natural 1557  
 curcuminoids. *Cancer Lett* 1995;94:79-83. 1558
- [135] Aggarwal BB, Shishodia S, Takada Y, Banerjee S, Newman 1559  
 RA, Bueso-Ramos CE, et al. Curcumin suppresses the 1560  
 paclitaxel-induced nuclear factor-kappaB pathway in 1561  
 breast cancer cells and inhibits lung metastasis of human 1562  
 breast cancer in nude mice. *Clin Cancer Res* 2005;11:7490- 1563  
 8. 1564
- [136] Bachmeier B, Nerlich AG, Iancu CM, Cilli M, Schleicher E, 1565  
 Vene R, et al. The chemopreventive polyphenol curcumin 1566  
 prevents hematogenous breast cancer metastases in 1567  
 immunodeficient mice. *Cell Physiol Biochem* 2007;19:137- 1568  
 52. 1569
- [137] Li L, Ahmed B, Mehta K, Kurzrock R. Liposomal curcumin 1570  
 with and without oxaliplatin: effects on cell growth, 1571  
 apoptosis, and angiogenesis in colorectal cancer. *Mol* 1572  
*Cancer Ther* 2007;6:1276-82. 1573
- [138] Cui SX, Qu XJ, Xie YY, Zhou L, Nakata M, Makuuchi M, 1574  
 et al. Curcumin inhibits telomerase activity in human 1575  
 cancer cell lines. *Int J Mol Med* 2006;18:227-31. 1576
- [139] Ohashi Y, Tsuchiya Y, Koizumi K, Sakurai H, Saiki I. 1577  
 Prevention of intrahepatic metastasis by curcumin in an 1578  
 orthotopic implantation model. *Oncology* 2003;65:250-8. 1579
- [140] LoTempio MM, Veena MS, Steele HL, Ramamurthy B, 1580  
 Ramalingam TS, Cohen AN, et al. Curcumin suppresses 1581  
 growth of head and neck squamous cell carcinoma. *Clin* 1582  
*Cancer Res* 2005;11:6994-7002. 1583
- [141] Odoj J, Albert P, Carlier A, Tarpin M, Devy J, Madoulet C. In 1584  
 vitro and in vivo anti-tumoral effect of curcumin against 1585  
 melanoma cells. *Int J Cancer* 2004;111:381-7. 1586
- [142] Lin YG, Kunnumakkara AB, Nair A, Merritt WM, Han LY, 1587  
 Arnaiz-Pena GN, et al. Curcumin inhibits tumor growth 1588  
 and angiogenesis in ovarian carcinoma by targeting the 1589  
 nuclear factor- $\kappa$ B pathway. *Clin Cancer Res* 1590  
 2007;13:3423-30. 1591
- [143] Li L, Braiteh FS, Kurzrock R. Liposome-encapsulated 1592  
 curcumin: in vitro and in vivo effects on proliferation, 1593  
 apoptosis, signaling, and angiogenesis. *Cancer* 1594  
 2005;104:1322-31. 1595
- [144] Kunnumakkara AB, Guha S, Krishnan S, Diagaradjane P, 1596  
 Gelovani J, Aggarwal BB. Curcumin potentiates antitumor 1597  
 activity of gemcitabine in an orthotopic model of 1598  
 pancreatic cancer through suppression of proliferation, 1599  
 angiogenesis, and inhibition of nuclear factor-kappaB- 1600  
 regulated gene products. *Cancer Res* 2007;67:3853-61. 1601
- [145] Dorai T, Cao YC, Dorai B, Buttyan R, Katz AE. Therapeutic 1602  
 potential of curcumin in human prostate cancer. III. 1603  
 Curcumin inhibits proliferation, induces apoptosis, and 1604  
 inhibits angiogenesis of LNCaP prostate cancer cells in 1605  
 vivo. *Prostate* 2001;47:293-303. 1606
- [146] Hong JH, Ahn KS, Bae E, Jeon SS, Choi HY. The effects of 1607  
 curcumin on the invasiveness of prostate cancer in vitro 1608  
 and in vivo. *Prostate Cancer Prostatic Dis* 2006;9:147-52. 1609
- [147] Menon LG, Kuttan R, Kuttan G. Inhibition of lung 1610  
 metastasis in mice induced by B16F10 melanoma cells by 1611  
 polyphenolic compounds. *Cancer Lett* 1995;95:221-5. 1612
- [148] Yoysungnoen P, Wirachwong P, Bhattarakosol P, Niimi H, 1613  
 Patumraj S. Effects of curcumin on tumor angiogenesis 1614  
 and biomarkers, COX-2 and VEGF, in hepatocellular 1615  
 carcinoma cell-implanted nude mice. *Clin Hemorheol* 1616  
*Microcirc* 2006;34:109-15. 1617
- [149] Busquets S, Carbo N, Almendro V, Quiles MT, Lopez- 1618  
 Soriano FJ, Argiles JM. Curcumin, a natural product 1619  
 present in turmeric, decreases tumor growth but does not 1620  
 behave as an anticachectic compound in a rat model. 1621  
*Cancer Lett* 2001;167:33-8. 1622
- [150] Wahlstrom B, Blennow G. A study on the fate of curcumin 1623  
 in the rat. *Acta Pharmacol Toxicol (Copenh)* 1978;43:86-92. 1624

- 1625 [151] Holder GM, Plummer JL, Ryan AJ. The metabolism and  
1626 excretion of curcumin (1,7-bis-(4-hydroxy-3-  
1627 methoxyphenyl)-1,6-heptadiene-3,5-dione) in the rat.  
1628 *Xenobiotica* 1978;8:761-8.
- 1629 [152] Ravindranath V, Chandrasekhara N. Absorption and  
1630 tissue distribution of curcumin in rats. *Toxicology*  
1631 1980;16:259-65.
- 1632 [153] Ravindranath V, Chandrasekhara N. Metabolism of  
1633 curcumin—studies with [3H]curcumin. *Toxicology*  
1634 1981;22:337-44.
- 1635 [154] Ireson C, Orr S, Jones DJ, Verschoyle R, Lim CK, Luo JL,  
1636 et al. Characterization of metabolites of the  
1637 chemopreventive agent curcumin in human and rat  
1638 hepatocytes and in the rat in vivo, and evaluation of their  
1639 ability to inhibit phorbol ester-induced prostaglandin E2  
1640 production. *Cancer Res* 2001;61:1058-64.
- 1641 [155] Ireson CR, Jones DJ, Orr S, Coughtrie MW, Boocock DJ,  
1642 Williams ML, et al. Metabolism of the cancer  
1643 chemopreventive agent curcumin in human and rat  
1644 intestine. *Cancer Epidemiol Biomarkers Prev* 2002;11:105-  
1645 11.
- 1646 [156] Garcea G, Jones DJ, Singh R, Dennison AR, Farmer PB,  
1647 Sharma RA, et al. Detection of curcumin and its  
1648 metabolites in hepatic tissue and portal blood of patients  
1649 following oral administration. *Br J Cancer* 2004;90:1011-5.
- 1650 [157] Hoehle SI, Pfeiffer E, Solyom AM, Metzler M. Metabolism of  
1651 curcuminoids in tissue slices and subcellular fractions  
1652 from rat liver. *J Agric Food Chem* 2006;54:756-64.
- 1653 [158] Ryu EK, Choe YS, Lee KH, Choi Y, Kim BT. Curcumin and  
1654 dehydrozingerone derivatives: synthesis, radiolabeling,  
1655 and evaluation for beta-amyloid plaque imaging. *J Med*  
1656 *Chem* 2006;49:6111-9.
- 1657 [159] Cheng AL, Hsu CH, Lin JK, Hsu MM, Ho YF, Shen TS, et al.  
1658 Phase I clinical trial of curcumin, a chemopreventive  
1659 agent, in patients with high-risk or pre-malignant lesions.  
1660 *Anticancer Res* 2001;21:2895-900.
- 1661 [160] Sharma RA, McLelland HR, Hill KA, Ireson CR, Euden SA,  
1662 Manson MM, et al. Pharmacodynamic and  
1663 pharmacokinetic study of oral *Curcuma* extract in patients  
1664 with colorectal cancer. *Clin Cancer Res* 2001;7:1894-900.
- 1665 [161] Garcea G, Berry DP, Jones DJ, Singh R, Dennison AR,  
1666 Farmer PB, et al. Consumption of the putative  
1667 chemopreventive agent curcumin by cancer patients:  
1668 assessment of curcumin levels in the colorectum and  
1669 their pharmacodynamic consequences. *Cancer Epidemiol*  
1670 *Biomarkers Prev* 2005;14:120-5.
- 1671 [162] Shoba G, Joy D, Joseph T, Majeed M, Rajendran R, Srinivas  
1672 PS. Influence of piperine on the pharmacokinetics of  
1673 curcumin in animals and human volunteers. *Planta Med*  
1674 1998;64:353-6.
- 1675 [163] Marczylo TH, Verschoyle RD, Cooke DN, Morazzoni P,  
1676 Steward WP, Gescher AJ. Comparison of systemic  
1677 availability of curcumin with that of curcumin formulated  
1678 with phosphatidylcholine. *Cancer Chemother Pharmacol*  
1679 2007;60:171-7.
- 1680 [164] Bisht S, Feldmann G, Soni S, Ravi R, Karikar C, Maitra A.  
1681 Polymeric nanoparticle-encapsulated curcumin  
1682 ("nanocurcumin"): a novel strategy for human cancer  
1683 therapy. *J Nanobiotechnol* 2007;5:3.
- 1684 [165] Sun A, Shoji M, Lu YJ, Liotta DC, Snyder JP. Synthesis of  
1685 EF24-tripeptide chloromethyl ketone: a novel curcumin-  
1686 related anticancer drug delivery system. *J Med Chem*  
1687 2006;49:3153-8.
- 1688 [166] Sandur SK, Pandey MK, Sung B, Ahn KS, Murakami A,  
1689 Sethi G, et al. Curcumin, demethoxycurcumin,  
1690 bisdemethoxycurcumin, tetrahydrocurcumin, and  
1691 turmerones differentially regulate anti-inflammatory and  
1692 antiproliferative responses through a ROS-independent  
1693 mechanism. *Carcinogenesis* 2007.
- [167] Pfeiffer E, Hoehle SI, Walch SG, Riess A, Solyom AM, Metzler M. Curcuminoids form reactive glucuronides in vitro. *J Agric Food Chem* 2007;55:538-44.
- [168] Sugiyama Y, Kawakishi S, Osawa T. Involvement of the beta-diketone moiety in the antioxidative mechanism of tetrahydrocurcumin. *Biochem Pharmacol* 1996;52:519-25.
- [169] Okada K, Wangpoengtrakul C, Tanaka T, Toyokuni S, Uchida K, Osawa T. Curcumin and especially tetrahydrocurcumin ameliorate oxidative stress-induced renal injury in mice. *J Nutr* 2001;131:2090-5.
- [170] Naito M, Wu X, Nomura H, Kodama M, Kato Y, Osawa T. The protective effects of tetrahydrocurcumin on oxidative stress in cholesterol-fed rabbits. *J Atheroscler Thromb* 2002;9:243-50.
- [171] Pari L, Amali DR. Protective role of tetrahydrocurcumin (THC) an active principle of turmeric on chloroquine induced hepatotoxicity in rats. *J Pharm Pharm Sci* 2005;8:115-23.
- [172] Murugan P, Pari L. Effect of tetrahydrocurcumin on plasma antioxidants in streptozotocin-nicotinamide experimental diabetes. *J Basic Clin Physiol Pharmacol* 2006;17:231-44.
- [173] Satoskar RR, Shah SJ, Shenoy SG. Evaluation of anti-inflammatory property of curcumin (diferuloyl methane) in patients with postoperative inflammation. *Int J Clin Pharmacol Ther Toxicol* 1986;24:651-4.
- [174] Kuttan R, Sudheeran PC, Joseph CD. Turmeric and curcumin as topical agents in cancer therapy. *Tumori* 1987;73:29-31.
- [175] Soni KB, Kuttan R. Effect of oral curcumin administration on serum peroxides and cholesterol levels in human volunteers. *Indian J Physiol Pharmacol* 1992;36:273-5.
- [176] Ramirez Bosca A, Soler A, Carrion-Gutierrez MA, Pamies Mira D, Pardo Zapata J, Diaz-Alperi J, et al. An hydroalcoholic extract of *Curcuma longa* lowers the abnormally high values of human-plasma fibrinogen. *Mech Ageing Dev* 2000;114:207-10.
- [177] James JS. Curcumin: clinical trial finds no antiviral effect. *AIDS Treat News* 1996;1-2.
- [178] Rasyid A, Lelo A. The effect of curcumin and placebo on human gall-bladder function: an ultrasound study. *Aliment Pharmacol Ther* 1999;13:245-9.
- [179] Rasyid A, Rahman AR, Jaalam K, Lelo A. Effect of different curcumin dosages on human gall bladder. *Asia Pac J Clin Nutr* 2002;11:314-8.
- [180] Lal B, Kapoor AK, Asthana OP, Agrawal PK, Prasad R, Kumar P, et al. Efficacy of curcumin in the management of chronic anterior uveitis. *Phytother Res* 1999;13:318-22.
- [181] Lal B, Kapoor AK, Agrawal PK, Asthana OP, Srimal RC. Role of curcumin in idiopathic inflammatory orbital pseudotumours. *Phytother Res* 2000;14:443-7.
- [182] Heng MC, Song MK, Harker J, Heng MK. Drug-induced suppression of phosphorylase kinase activity correlates with resolution of psoriasis as assessed by clinical, histological and immunohistochemical parameters. *Br J Dermatol* 2000;143:937-49.
- [183] Sharma RA, Euden SA, Platton SL, Cooke DN, Shafayat A, Hewitt HR, et al. Phase I clinical trial of oral curcumin: biomarkers of systemic activity and compliance. *Clin Cancer Res* 2004;10:6847-54.
- [184] Bundy R, Walker AF, Middleton RW, Booth J. Turmeric extract may improve irritable bowel syndrome symptomology in otherwise healthy adults: a pilot study. *J Altern Complement Med* 2004;10:1015-8.
- [185] Shoskes D, Lapiere C, Cruz-Correa M, Muruve N, Rosario R, Fromkin B, et al. Beneficial effects of the bioflavonoids curcumin and quercetin on early function in cadaveric renal transplantation: a randomized placebo controlled trial. *Transplantation* 2005;80:1556-9.

- 1763 [186] Durgaprasad S, Pai CG, Vasanthkumar, Alvres JF, Namitha S. A pilot study of the antioxidant effect of curcumin in  
1764 tropical pancreatitis. *Indian J Med Res* 2005;122:315–8.  
1765 [187] Holt PR, Katz S, Kirshoff R. Curcumin therapy in  
1766 inflammatory bowel disease: a pilot study. *Dig Dis Sci*  
1767 2005;50:2191–3.  
1768 [188] Hanai H, Iida T, Takeuchi K, Watanabe F, Maruyama Y,  
1769 Andoh A, et al. Curcumin maintenance therapy for  
1770 ulcerative colitis: randomized, multicenter, double-blind,  
1771 placebo-controlled trial. *Clin Gastroenterol Hepatol*  
1772 2006;4:1502–6.  
1773 [189] Cruz-Correa M, Shoskes DA, Sanchez P, Zhao R, Hyland  
1774 LM, Wexner SD, et al. Combination treatment with  
1775 curcumin and quercetin of adenomas in familial  
1776 adenomatous polyposis. *Clin Gastroenterol Hepatol*  
1777 2006;4:1035–8.  
1778 [190] Ng TP, Chiam PC, Lee T, Chua HC, Lim L, Kua EH. Curry  
1779 consumption and cognitive function in the elderly. *Am J*  
1780 *Epidemiol* 2006;164:898–906.  
1781 [191] Rafailov S, Cammack S, Stone BA, Katz AE. The role of  
1782 zyflamend, an herbal anti-inflammatory, as a potential  
1783 chemopreventive agent against prostate cancer: a case  
1784 report. *Integr Cancer Ther* 2007;6:74–6.  
1785 [192] Di Mario F, Cavallaro LG, Nouvenne A, Stefani N, Cavestro  
1786 GM, Iori V, et al. A curcumin-based 1-week triple therapy  
1787 for eradication of *Helicobacter pylori* infection: something to  
1788 learn from failure? *Helicobacter* 2007;12:238–43.  
1789 [193] Lao CD, Demierre MF, Sondak VK. Targeting events in  
1790 melanoma carcinogenesis for the prevention of  
1791 melanoma. *Expert Rev Anticancer Ther* 2006;6:1559–68.  
1792 [194] Kobelt G. Health economic issues in rheumatoid arthritis.  
1793 *Scand J Rheumatol* 2006;35:415–25.  
1794 [195] Chen YC, Tsai SH, Shen SC, Lin JK, Lee WR. Alternative  
1795 activation of extracellular signal-regulated protein  
1796 kinases in curcumin and arsenite-induced HSP70 gene  
1797 expression in human colorectal carcinoma cells. *Eur J Cell*  
1798 *Biol* 2001;80:213–21.  
1800 [196] Dhillon N, Wolff RA, Abbruzzese JL, et al. Phase II clinical  
1801 trial of curcumin in patients with advanced pancreatic  
1802 cancer. *J Clin Oncol* 2006;24:14151 [abstract].
- [197] Frank N, Knauff J, Amelung F, Nair J, Wesch H, Bartsch H. 1803  
No prevention of liver and kidney tumors in Long-Evans 1804  
Cinnamon rats by dietary curcumin, but inhibition at 1805  
other sites and of metastases. *Mutat Res* 2003;523– 1806  
524:127–35. 1807
- [198] Aggarwal BB, Banerjee S, Bharadwaj U, Sung B, Shishodia 1808  
S, Sethi G. Curcumin induces the degradation of cyclin E 1809  
expression through ubiquitin-dependent pathway and up- 1810  
regulates cyclin-dependent kinase inhibitors p21 and p27 1811  
in multiple human tumor cell lines. *Biochem Pharmacol* 1812  
2007;73:1024–32. 1813
- [199] Moos PJ, Edes K, Mullally JE, Fitzpatrick FA. Curcumin 1814  
impairs tumor suppressor p53 function in colon cancer 1815  
cells. *Carcinogenesis* 2004;25:1611–7. 1816
- [200] Tsvetkov P, Asher G, Reiss V, Shaul Y, Sachs L, Lotem J. 1817  
Inhibition of NAD(P)H:quinone oxidoreductase 1 activity 1818  
and induction of p53 degradation by the natural phenolic 1819  
compound curcumin. *Proc Natl Acad Sci USA* 1820  
2005;102:5535–40. 1821
- [201] Somasundaram S, Edmund NA, Moore DT, Small GW, Shi 1822  
YY, Orłowski JZ. Dietary curcumin inhibits 1823  
chemotherapy-induced apoptosis in models of human 1824  
breast cancer. *Cancer Res* 2002;62:3868–75. 1825
- [202] Hata M, Sasaki E, Ota M, Fujimoto K, Yajima J, Shichida T, 1826  
et al. Allergic contact dermatitis from curcumin 1827  
(turmeric). *Contact Dermat* 1997;36:107–8. 1828
- [203] Swjerczynska MK, Krecisz B. Occupational skin changes in 1829  
persons working in contact with food spices. *Med Pr* 1830  
1998;49:187–90. 1831
- [204] Keijth CT, Borisy AA, Stockwell BR. Multicomponent 1832  
therapeutics for networked systems. *Nat Rev Drug Discov* 1833  
2005;4:71–8. 1834
- [205] Sams-Dodd F. Target-based drug discovery: is something 1835  
wrong? *Drug Discov Today* 2005;10:139–47. 1836
- [206] Morphy R, Kay C, Rankovic Z. From magic bullets to 1837  
designed multiple ligands. *Drug Discov Today* 2004;9: 1838  
641–51. 1839
- [207] Mencher SK, Wang LG. Promiscuous drugs compared to 1840  
selective drugs (promiscuity can be a virtue). *BMC Clin* 1841  
*Pharmacol* 2005;5:3. 1842  
1843