Phytochemicals as Modulators of Neoplastic Phenotypes

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Introduction

The value of nonessential, natural molecules contained in fruits and vegetables (micronutrients or phytochemicals) in maintaining health has steadily gained acceptance over the past two decades. The results of epidemiological studies on the potential benefits of fruit and vegetable diets [1], and molecular studies on the cellular effects of an extensive list of phytochemicals [2, 3], have generated the general awareness that molecules of vegetal origin are effective and beneficial modulators of cellular function and contribute to the prevention of cancer [4].

Representative classes of dietary bioactive molecules including terpenoids, polyphenolics, organosulfurous and alkaloid compounds are presented in table 1.

Key Words
Phytochemicals · Cancer prevention · Oncogenic addiction · Hormesis · Membrane material properties · Genome instability

Abstract

It is generally accepted that nutritional behaviors constitute decisive components of human health. Phytochemicals (small, nonenergetic molecules of vegetal origin) are overall inhibitory on the expression of gene products promoting proliferation and resistance to apoptosis. On the contrary, phytochemicals stimulate the synthesis of adaptive proteins that favor resistance to cellular stress (detoxifying and antioxidant enzymes). They are effective modulators that act synergistically on membrane, cytoplasmic and nuclear enzymatic reactions to dampen cellular hyperproliferation and hyperactivity, reequilibrate metabolic activity and promote apoptosis of genetically unstable cells. Despite important gaps in our knowledge regarding how phytochemicals interfere with cellular function in vivo, effective chemopreventive measures have shown that phytochemicals can be utilized to prevent cancer, and possibly to treat cancer patients as well. We review how phytochemicals exert their beneficial effects at the cellular level.
teins. In vitro studies have identified the signaling pathways modulated by phytochemicals that regulate the activity of major transcription factors such as NFκB (nuclear factor-kappa B), FOXO (forkhead box group O) and Nrf2 (NF-E2-related factor-2), that respectively control antioxidant responses, detoxifying enzyme synthesis, energy metabolism and survival [7].

Given such effects on basic life-saving and adaptive functions, how then could phytochemicals exert an anti-cancer effect and selectively bring about the demise of neoplastic cells? Phytochemicals characteristically affect fundamental cellular signaling pathways by interfering with many protein or lipid kinases and transcription factors. They are essentially pleiotropic and able to inter-

<table>
<thead>
<tr>
<th>Phytochemical</th>
<th>Plant</th>
<th>Molecular structure</th>
</tr>
</thead>
<tbody>
<tr>
<td>Curcumin</td>
<td>turmeric</td>
<td><img src="image_url" alt="Curcumin" /></td>
</tr>
<tr>
<td>Resveratrol</td>
<td>(red) grapes, berries, peanuts, rhubarb</td>
<td><img src="image_url" alt="Resveratrol" /></td>
</tr>
<tr>
<td>Epicatechin</td>
<td>cocoa, grapes, tea, apples</td>
<td><img src="image_url" alt="Epicatechin" /></td>
</tr>
<tr>
<td>Genistein</td>
<td>soybean, (chickpeas)</td>
<td><img src="image_url" alt="Genistein" /></td>
</tr>
<tr>
<td>Quercetin</td>
<td>onions, apples, tea, nuts, berries, cauliflower, cabbage</td>
<td><img src="image_url" alt="Quercetin" /></td>
</tr>
<tr>
<td>Caffeic acid</td>
<td>pears, basil, thyme, verbena, oregano, rosemary, coffee</td>
<td><img src="image_url" alt="Caffeic acid" /></td>
</tr>
<tr>
<td>Capsaicin</td>
<td>red chillies</td>
<td><img src="image_url" alt="Capsaicin" /></td>
</tr>
<tr>
<td>Lycopene</td>
<td>tomatoes, watermelons, guavas, pink grapefruits, red oranges</td>
<td><img src="image_url" alt="Lycopene" /></td>
</tr>
<tr>
<td>β-Carotene</td>
<td>mangos, apricots, carrots, broccoli, spinach</td>
<td><img src="image_url" alt="β-Carotene" /></td>
</tr>
<tr>
<td>Sulforaphane</td>
<td>broccoli, cauliflower, cabbage, kale</td>
<td><img src="image_url" alt="Sulforaphane" /></td>
</tr>
<tr>
<td>Allicin</td>
<td>garlic</td>
<td><img src="image_url" alt="Allicin" /></td>
</tr>
</tbody>
</table>

*Terpenoids: carotenoid terpenoids, e.g. lycopene, β-carotene; alkaloid: e.g. capsaicin; polyphenolics: flavonoid polyphenolics, e.g. catechins, quercetin, genistein; phenolic acids, e.g. caffeic acid; other nonflavonoid polyphenolics, e.g. curcumin, resveratrol; organosulfur compounds: isothiocyanates, e.g. sulforaphane; thiosulfonates, e.g. allicin.*
fere with pathways at different levels, not only by blocking enzymes or transcription factor binding to specific DNA response elements, but also by interfering with the membrane structure, intracytoplasmic protein-protein interactions, catalytic function of enzymes and last but not least, by interacting with DNA and promoting genome stability and integrity. As a result of such ‘nonspecific’ interferences, phytochemicals disorganize the aberrant signaling of neoplastic cells and may restore their apoptotic potential [4], as they are able to restore a normal activity profile in chronically active inflammatory cells [8]. In addition to their useful inhibitory effects on isolated enzymes (for instance, quercetin [9] and resveratrol [10] on PI3K (phosphatidylinositol 3’-kinase), and genistein on Src kinases [11], or piceatannol, a metabolite of resveratrol, on the Syk kinase [12]), phytochemicals like resveratrol exert their pleiotropic inhibitory effects with comparable success on neoplastic cells of different origins that most probably utilize different signaling circuits. A key survival transcription factor, such as STAT3 (signal transducer and activator of transcription-3) was efficiently inhibited by resveratrol [13] and ursolic acid [14].

One possible cause for the increased susceptibility of neoplastic and possibly other non-neoplastic (inflammatory) cells could be the signaling specialization (‘oncogenic addiction’) observed in epithelial [15, 16] and hematopoietic [17] cancer cells for one or a few oncogenic signaling proteins. With such dysregulated signaling networks strongly biased towards a limited set of signaling proteins, it is indeed understandable that phytochemicals could block prosurvival pathways more easily than in a non-neoplastic cell relying on several pathways to carry out a comparable task. Phytochemicals also influence carcinogenesis by strongly increasing the endogenous antioxidant potential [18], an enhanced capacity which is probably very effective in situations where chronic inflammation generates oxidants that pave the way for carcinogenesis [19].

The goal of this review is to discuss the anticancer effects of phytochemicals at the critical locations where signaling can be interfered with, namely (1) the plasma membrane as the site of proximal signaling, (2) the cytoplasmic signaling steps at which signalosomes are assembled and transcription factors activated, and (3) the nucleus, where activated transcription factors recognize their DNA response elements. To conclude, the important issue of how to exploit phytochemicals shall be discussed, in particular the issue of bioavailability. Most of the studies presently available have investigated in vitro how phytochemicals interact with target cells. Although such studies provide invaluable information about how phytochemicals bind and modulate specific enzymes and how they chemically modify proteins and modulate redox signaling, it should be kept in mind that much remains to be learned about how phytochemicals exert their effects in vivo. Although many mouse model studies [20] and preclinical studies [21] have provided ample proof of in vivo efficacy, exactly which chemical forms of the plant molecule act on the cancer cell target and how this affects cellular functions remains to be defined.

### How Phytochemicals Interact with Target Cells

The cell-modulating effects of phytochemicals shown for different cellular compartments and discussed on the following pages are summarized in figure 1.

#### Plasma Membrane Effects

The plasma membrane constitutes the first cellular organelle coming in contact with extracellular agonists, antagonists and toxic molecules, and contains receptors and channel proteins capable of recognizing and importing such molecules. Hydrophobic extracellular molecules could also directly interact with the membrane bilayer, and thus alter the receptors and channels that allosterically depend on their membrane lipid milieu for function [22]. This dependence of receptors and their associated cytoplasmic signaling effectors on the membrane lipids assembled in a microdomain (raft) has already been included in the concept of raft-associated signaling platforms, according to which functional signaling complexes preferentially form and function in sphingolipid-enriched and liquid-ordered plasma membrane microdomains [23, 24]. An abundant literature has been devoted to the characterization of the interaction of phytochemicals with plasma membranes, and we shall highlight their most relevant effects on cancer cell membrane structure and function.

#### Modulation of Membrane Material Properties

Curcumin (diferuloylmethane) is a major bioactive compound in the turmeric spice (Curcuma longa) that especially accumulates in tumor cell membranes [25]. This accumulation in membranes causes alterations of cell shape and modulates the bilayer material properties (bilayer thickness, fluidity and elasticity) that affect
membrane proteins (for instance, epidermal growth factor receptor (EGFR), multidrug resistance proteins, cystic fibrosis transmembrane conductance receptor (CFTR) and Kv1.4 potassium channels [26]). Curcumin does not target any membrane protein in particular, but rather modifies the hydrophobic coupling between membrane proteins and neighboring lipids in the membrane. In artificial bilayers, curcumin was shown to affect gramicidin single channel lifetime by causing a nonlinear membrane thinning [27]. Very similar effects have been reported for capsaicin [28] and genistein [29], and the liposomal membrane fluidity was altered in the same manner by flavonoid polyphenolics [30].

The lipophilic lycopenes and β-carotenes as well as *Allium* organosulfurous compounds also alter the membrane material properties (for instance, by rigidification of the hydrophobic core of the bilayer caused by denser packing of phospholipid acyl chains and increased membrane width [31, 32]). The amphiphilic resveratrol increases membrane fluidity and may also induce relocation of the Fas receptors to rafts via interaction of its polar groups with the amino groups of the transmembrane Fas receptors [33, 34].

**Perturbation of Enzymes Associated with the Inner Membrane Leaflet**

Critical signaling proteins such as Src family kinases [35] or PI3K/Akt kinases [36] are associated with the cytoplasmic leaflet of the plasma membrane. Genistein is a Src family kinase inhibitor [11] that also modulates chan-
nel function in a phosphorylation-independent manner, probably by dissociating Src family kinases from their optimal membrane lipid environment through rigidifying the cytoplasmic leaflet [29]. It is therefore expected that PI3K/Akt and protein kinase C (PKC) associations with the cytoplasmic membrane leaflet will be affected by phytochemicals that alter the physical properties of the plasma membrane, and particularly those of the inner leaflet. It could be envisaged that quercetin, by intercalating in the hydrophobic core of the bilayer, could stabilize the outer leaflet lipids in a manner analogous to cholesterol when it is embedded in sphingolipid-rich domains and liquid-ordered raft microdomains [37]. Likewise, flavonoids incorporated into erythrocyte ghost membranes increase lipid order with concomitant increases in antioxidant and antihemolytic properties [38].

Resveratrol inserts into the hydrophobic membrane core, causing for instance an inhibition of PKC association with the inner leaflet [39]. Inhibition of other inner leaflet-associated signaling kinases such as PI3K [10] and Akt are also consequences of in vitro exposure to resveratrol [40].

Effects of Phytochemicals on Membrane Lipid Oxidation

Polyphenolics are generally amphiphilic and their degree of hydrophobicity or hydrophilicity depends on the number of polar head groups. The surface-bound polyphenolics may stabilize the bilayer [41], but pH influences their hydrophobicity with the effect that polyphenolics of increased hydrophobicity intercalate in the lipid leaflets. For instance, quercetin was shown to become embedded in the lipid bilayers at acidic pH, but to intercalate in the hydrophobic core of the bilayer, could stabilize the outer leaflet lipids in a manner analogous to cholesterol when it is embedded in sphingolipid-rich domains and liquid-ordered raft microdomains [37]. Likewise, flavonoids incorporated into erythrocyte ghost membranes increase lipid order with concomitant increases in antioxidant and antihemolytic properties [38].

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For curcumin, resveratrol, epigallocatechin gallate, caffeic acid phenetyl ester (CAPE) and isothiocyanates, inhibition of Nrf2 and AP-1 transcription factors was reported while the nuclear translocation of Nrf2, its binding to antioxidant response elements (AREs) and the expression of detoxifying enzymes were increased [45]. For selected phytochemicals bearing an unsaturated keto moiety (i.e. curcumin, CAPE and sulforaphane), a direct interaction of those ketone moieties with thiol groups of the Keap1 (kelch-like ECH-associated protein 1) protein has been reported. The Keap1 protein that forms a complex with Nrf2 and sequesters it in the cytoplasm [46], can be modified by phytochemicals to release Nrf2 and allow its interaction with antioxidant response elements in the nucleus to enhance the expression of phase II detoxifying and antioxidant enzymes [47]. Phase I enzymes hydrolyze, reduce or oxidize substrates which are further processed by phase II transferases conjugating glucuronate, sulfate or amino acids to the metabolites to augment their water solubility.

Much of the biochemical information regarding the way in which phytochemicals inhibit kinases is still lacking, but many natural products do compete with ATP for the active site of kinases. For instance, resveratrol targets the ATP-binding site of class IA PI3K in a competitive and reversible fashion, independently of its activation of SIRT1 (sirtuin1, silent mating type information regula-
The maintenance of genome integrity is achieved by an elaborate machinery of DNA repair enzymes that excise damaged DNA and synthesize new DNA to ensure the availability of undamaged genetic material [53]. The usefulness of several phytochemicals in maintaining genome integrity was recently reviewed [54] and shows that phytochemicals do far more than slowing down the process of carcinogenesis, as originally documented for resveratrol [55]. A direct interaction of resveratrol, genistein and quercetin with DNA [56, 57] indeed suggests that such phytochemicals may exert a protective effect directly on DNA by preventing breaks [54].

The ATM/ATR (ataxia telangiectasia mutated kinase/ataxia telangiectasia and Rad3-related kinase) kinases are PI3K-related kinases involved in double-strand break repair, and activated by resveratrol [58]. However, there remain many unknown factors regarding the causes of ATM/ATR activation by resveratrol. Interestingly, resveratrol (but also quercetin and genistein) appear capable of activating the ATM/ATR kinases without previous DNA damage, and thus may prepare the cell to face further challenges such as those encountered in the course of cancer progression. This is indeed in line with the hormetic effects attributed to many phytochemicals [7]. Despite a number of contradictory observations, it seems probable that SIRT1 is involved in resveratrol-mediated DNA repair via the SIRT1 deacetylated substrates p53 and Nbs1 [43].

Bioavailability

The question of how, in which chemical form and to what extent phytochemicals gain access to cells and tissues is central to understanding their mode of action. The majority of phytochemicals exist in conjugated form and only a small amount of phytochemicals are present in free form. The modifications (glucuronidation and sulfation by phase II enzymes) lower the bioavailability of the substances by reducing their cell permeability or absorption [59]. Conjugated phytochemicals that are not absorbed in the upper digestive tract may be broken down into smaller molecular weight molecules (for instance cinnamic acid) by the colonic microflora and display a high degree of bioavailability [60]. This aspect of the metabolism of phytochemicals has not been thoroughly investigated and holds a high potential for discovery [61].

It also remains to be investigated whether insoluble metabolites of phytochemicals constitute a spare pool of free active phytochemicals, or only inactive and excretable forms [20]. The influence of the food matrix on the bioavailability of phytochemicals likewise deserves further attention. Flavonoids which are mainly present as glycosides in food (with the exception of catechins) are expected to be poorly absorbed, but quercetin glycosides are absorbed in appreciable amounts in the small intestine [62]. For instance, the flavonoid quercetin was shown to be more bioavailable as an aglycone than quercetin glycosides when ingested as onion flesh, while quercetin glycosides where more available when ingested as dried onion skin [63]. The purified and hydrophobic quercetin aglycone, however, is almost not absorbed. There are also...
contradictory reports as to whether curcumin metabolites are more or less active when conjugated [64].

The maximal concentrations of conjugated and unconjugated resveratrol in the blood can be expected to reach 1–10 μM [65], concentrations at which phytochemicals may exert antiproliferative and apoptosis-promoting effects [18], Intestinal mucosal cells, however, may be exposed to much higher phytochemical concentrations than those utilized in vitro (up to 100–500 μM). To induce antioxidant and anticancer effects in vitro, experiments using single phytochemical application always require higher doses than those usually achieved in vivo under physiological conditions. However, the proven in vivo beneficial effects of phytochemicals in cancer prevention and suppression can be explained by additive and synergistic effects, as vegetables and fruits contain multiple different phytochemicals which seem to influence and potentiate each other [59–61]. Synergistic effects increasing the bioavailability are reported. For example quercetin is an inhibitor of resveratrol sulfation in the liver and small intestine and increases the bioavailability of resveratrol [20]. The synergistic effect of piperine on curcumin is driven by its inhibiting effect on curcumin conjugation. Further, absorption of phytochemicals can be enhanced by complexing with lipids or by nanoparticles that increase the water solubility of hydrophobic drugs [64].

Conclusions

The anticancer effects of phytochemicals manifest themselves both on the cancer cell itself, and on the initiated cell (a cell on its way to become neoplastic). The many pathways that are modified by phytochemicals slow down the inappropriately proliferating cells, most likely by participating in a generic mechanism of cellular regulation which is influenced by the cellular redox state.

As suggested by Stevenson and Hurst [18], humans have had a long time to adjust to the intrinsically noxious effects of plant phytochemicals and consequently have developed adaptive mechanisms to neutralize those phytochemicals. It is quite fascinating to realize that chimpanzees, the evolutionary cousins of man [66], have been shown to make use of plants to try to alleviate ailments [67, 68] in a manner analogous to what early humans must have done. Phytochemicals are therefore molecules that have shaped the adaptive and hormetic mechanisms in human physiology and have provided us with the capacity to regulate cellular functions.

The chemopreventive properties of phytochemicals appear sufficient to safely decrease the incidence of cancers due to environmental causes. Whether or not purified phytochemicals could be utilized as drugs to regulate aberrant cellular behaviors will be learned from the many clinical studies presently under way, and, hopefully, selected phytochemicals will find their use in our pharmacology to act in a pleiotropic manner on neoplastic, inflammatory and metabolically unbalanced cells. This could in essence be a new way to influence aberrant cellular behavior, not by seeking the elimination of one faulty molecule (the magic bullet approach), but rather by attempting to reequilibrate an unbalanced cellular machinery, such as in diabetic or cancer cells.

Acknowledgments

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References


Bloom DA, Jaiswal AK: Phosphorylation of Nrf2 at Ser40 by protein kinase C in response to antioxidants leads to the release of Nrf2 from IкBк, but is not required for Nrf2 stabilization/acclimation in the nucleus and transcriptional activation of antioxidant response element-mediated NAD(P)H:quinone oxidoreductase-1 gene expression. J Biol Chem 2003;278:44675–44682.