Pilot Study of Oral Silibinin, a Putative Chemopreventive Agent, in Colorectal Cancer Patients: Silibinin Levels in Plasma, Colorectum, and Liver and Their Pharmacodynamic Consequences

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Abstract

Silibinin, a flavonolignan from milk thistle, has intestinal cancer chemopreventive efficacy in rodents. It is a strong antioxidant and modulates the insulin-like growth factor (IGF) system by increasing circulating levels of IGF-binding protein 3 (IGFBP-3) and decreasing levels of IGF-1. Here, the hypothesis was tested that administration of oral silibinin generates agents levels in human blood and colorectal and hepatic tissues consistent with pharmacologic activity. Patients with confirmed colorectal adenocarcinoma received silibinin formulated with phosphatidylcholine (silipide) at dosages of 360, 720, or 1,440 mg silibinin daily for 7 days. Blood and biopsy samples of normal and malignant colorectum or liver were obtained before dosing, and blood and colorectal or hepatic tissues were collected at resection surgery after the final silipide dose. Levels of silibinin were quantified by high-pressure liquid chromatography-UV, and plasma metabolites were identified by liquid chromatography-mass spectrometry. Blood levels of IGFBP-3, IGF-I, and the oxidative DNA damage pyrimidopurinone adduct of deoxyguanosine (M1dG) were determined. Repeated administration of silipide was safe and achieved levels of silibinin of 0.3 to 4 μmol/L in the plasma, 0.3 to 2.5 nmol/g tissue in the liver, and 20 to 141 nmol/g tissue in colorectal tissue. Silibinin monoglucuronide, silibinin diglucuronide, silibinin monosulfate, and silibinin glucuronide sulfate were identified in the plasma. Intervention with silipide did not affect circulating levels of IGFBP-3, IGF-I, or M1dG. The high silibinin levels achieved in the human colorectal mucosa after consumption of safe silibinin doses support its further exploration as a potential human colorectal cancer chemopreventive agent.

Silibinin, a flavonolignan (for structure, see Fig. 1), is a major constituent of the seeds of milk thistle (Silybum marianum L.). Extracts of milk thistle are widely consumed as a dietary supplement especially in the United States. Silymarin, a standardized milk thistle extract, of which silibinin is the main component, has been evaluated clinically in the treatment of hepatitis and liver damage inflicted by alcohol and long-term treatment with psychotropic drugs (1–4). Recent evidence in rodents suggests that silymarin and silibinin may be useful in the chemoprevention of malignancies at a variety of sites, including the intestinal tract (5–10). Dietary silymarin delayed the development of intestinal adenocarcinomas in rats induced by dimethylhydrazine (8) or azoxymethane (9). It also suppressed aberrant crypt foci in rats, which had been exposed to azoxymethane, but this suppression was dose independent and did not involve induction of apoptosis (10). In our laboratory, silibinin interfered moderately with small intestinal carcinogenesis in the ApcMin mouse model. Silibinin has been formulated with phosphatidylcholine (silipide, Indena SpA, Milan, Italy) to improve its systemic availability, and clinical evaluation of this formulation at single or repeated doses reflects both its safety (summarized in ref. 11) and its improved bioavailability with respect to silymarin (12). Several mechanisms have been proposed to explain how silibinin may interfere with carcinogenesis. Among these mechanisms are impairment of receptor tyrosine kinase and erbB1 signaling and up-regulation of cyclin-dependent kinase inhibitors causing attenuation of cancer cell growth and perturbation of cell cycle progression (13, 14), induction of cancer cell differentiation (15), and antiangiogenesis (16). Silibinin has also been suggested to modulate the insulin-like growth factor (IGF) system. IGFs are mediators of cell survival in that they can inhibit apoptosis...
and influence differentiation of many normal and malignant cell types (17–19). The IGF system is regulated by IGF-binding proteins (IGFBP), especially IGFBP-3, which bind IGFs in the extracellular milieu with high affinity and specificity, thus reducing the bioavailability of IGFs. Epidemiologic studies have linked increased serum concentrations of IGF-I, decreased concentrations of IGFBP-3, or both with an increased risk of cancer at several sites, including the colorectum (20, 21). Silibinin increased levels of IGFBP-3 in prostate cells in vitro (22) and in tumor-bearing rodents in vivo (23). Silymarin and silibinin are also powerful antioxidants as a consequence of their polyphenolic structure (11, 24). Elevated oxidative stress has been implicated as an important carcinogenic stimulus, and endogenous or exogenous oxidative events, such as those associated with lipid peroxidation, generate malondialdehyde, a mutagen (25). Malondialdehyde can react with DNA to form the pyrimidopurinone adduct of deoxyguanosine M1dG (26). M1dG levels may be a suitable marker of chemopreventive efficacy of strong antioxidants, such as silibinin.

Silibinin undergoes extensive metabolism, especially phase II conjugation reactions, both in human liver preparations in vitro (27) and in volunteers in vivo after consumption of silibinin (28). There is no information available on the pharmacologic properties of metabolic conjugates of silibinin, and it is conceivable that at least some of them share pharmacologic activity with the parent molecule. Taken together, all of these results intimate both the propitiousness and the feasibility of developing silibinin as a colorectal cancer chemopreventive agent. To obtain pharmacokinetic and pharmacodynamic information to aid the design of future colorectal cancer intervention studies with silibinin, we carried out a pilot “presurgery model” study in colorectal cancer patients. Patients consumed silibinin daily for a week before colectomy, and blood and colorectal or hepatic tissue was obtained by biopsy and resection before and after the administration. Blood and tissues were analyzed to explore whether metabolites of silibinin can be identified in human blood and whether silibinin levels achievable in blood, colorectal, or hepatic tissue are comparable with pharmacologically active levels reported in cells in vitro. Furthermore we wished to find out if consumption of silibinin affects circulating levels of IGFBP-3, IGF-I, and the oxidative DNA damage adduct M1dG. Overall, the study was designed to provide data, which helps optimize the design of long-term intervention studies of silibinin as a colorectal cancer chemopreventive agent.

**Materials and Methods**

**Patients and tissues.** Twelve patients (1 female and 11 male, ages 55-78 years, mean age 65 ± 7) with confirmed colorectal carcinoma of stages A (2 patients), B (5), or C (5), who were to undergo colorectal resection, and 12 patients (7 female and 5 male, ages 49-78 years, mean age 62 ± 9, all Dukes D) with hepatic metastatic disease originating from primary colorectal carcinoma, who were to undergo hepatic surgery, were recruited into the trial following approval by the University Hospitals of Leicester (Leicester, United Kingdom) ethics committee. All patients gave written informed consent. Hematologic profiles, plasma levels of urea and electrolytes, and hepatic function were within the reference range defined by the laboratories of the University Hospitals of Leicester. One patient who underwent colectomy had preoperative radiotherapy and none preoperative chemotherapy. All, except two patients who underwent hepatic surgery, had received 5-fluorouracil with folinic acid, oxaliplatin, and/or irinotecan before recruitment. Drug histories included antihypertensive, diuretics, antidepressants, and analgesics. Colorectal tumors in patients who underwent colorectal resection were located in the ascending/transverse colon (one patient), sigmoid colon (four patients), or rectum (seven patients). Tissue biopsy specimens taken at diagnosis weighed 3 to 60 mg (tumor biopsy) and 4 to 60 mg (normal tissue biopsy). Colectomy or hepatic resection was done 3 to 6 hours after the last dose of silipide. The weight of the surgical colorectal/liver tissue samples for chemical analysis was 1.1 to 1.5 g. Samples of peripheral blood were collected in heparinized tubes before intervention and 1 to 4 hours after the last silipide dose. Portal blood was taken at the point of hepatic surgery. Blood samples were centrifuged to generate leukocytes (for M1dG measurement), serum (for IGF-I/IGFBP-3 measurement), and plasma [for high-pressure liquid chromatography (HPLC) analysis]. Blood components were kept on ice until storage at −80°C. Tissue samples were flash frozen in liquid nitrogen. Biomarkers were kept at −80°C for up to 6 months until analysis. Preliminary HPLC analysis had established that silibinin is stable under these conditions in tissues and plasma.

**Silibinin formulation and dose.** Silibinin was formulated in capsules as silipide (IdB 1016), a phytosome product marketed by Indena SpA for use as a hepatoprotectant (see www.indena.it/pdf/prodottiw eb.pdf). Each silipide capsule contained 120 mg silibinin (validated by HPLC-UV analysis, see below) and 30% phosphatidylcholine. Patients received silipide at dosages for use as a hepatoprotectant (see www.indena.it/pdf/prodottiweb.pdf). Each silipide capsule contained 120 mg silibinin (validated by HPLC-UV analysis, see below) and 30% phosphatidylcholine. Patients received silipide at dosages of either 360, 720, or 1,440 mg silibinin daily for 7 days before surgery; each daily dose was divided in three equal portions taken at intervals in the morning, at noon, and in the evening. There were eight individuals per dose level (four patients who underwent colectomy and four who had liver resection). The first and second portions of the first dose were taken at noon and in the evening, respectively, of day 1; the last dose portion was ingested in the morning of day 8 before surgery so that, in total, the seven daily doses were distributed ×8 days. The doses used in this pilot study were based approximately on the dose that raised plasma IGFBP-3 levels in mice (23). In that study, mice received silibinin at 0.05% or 0.1% in the diet, which equates to ~75 and 150 mg/kg daily, respectively. These doses extrapolated to humans based on body surface area (420 mg/m² in mice; ref. 29) would amount to ~450 or ~900 mg silibinin/person daily, assuming a body surface area of 1.8 m² accompanying a body weight of 70 kg. Although such extrapolation has to be interpreted with caution, the doses proposed for the pilot study here cover the dose shown to be efficacious in mice.

**Analysis of silibinin and silibinin conjugates.** Biomarkers were thawed, weighed, and homogenized in an equal part of KCl solution (0.15 mol/L). Samples of plasma or tissue homogenate were mixed with 3 parts ice-cold methanol. The mixture was centrifuged, and the supernatant was decanted and dried under nitrogen, reconstituted in aqueous methanol (70%, containing 5% acetic acid), and analyzed for silibinin by HPLC with UV detection using a gradient system with a
two-component mobile phase. The details, characterization, and validation of the method are described in a separate article. The limit of quantitation for the two silibinin diastereoisomers was 5 and 3 ng/mL. Some plasma samples were incubated (1 hour, 37°C, 0.1 mol/L ammonium acetate buffer (pH 6.0) with β-glucuronidase from Helix pomatia (type H-2, containing sulfatase; Sigma-Aldrich, Poole, United Kingdom) before extraction.

Silibinin and some of its metabolites were characterized by LC-mass spectrometry (MS) using a turbospray source in negative electrospray ionization mode. Analyses were done using an API 2000 LC-MS (Applied Biosystems/MDS Sciex, Warringon, United Kingdom) equipped with an Agilent (South Queensferry, United Kingdom) 1100 series sample delivery system. Separation of silibinin and metabolites was achieved using the method mentioned above. MS conditions were as follows: declustering potential, −26 V; focusing potential, −350 V; electrode potential, −12 V; cell entrance potential, −16 V; cell exit potential, −20 V; and temperature, 500°C. Identification of silibinin-derived species was by selected ion monitoring.

Pharmacodynamic measurements. IGF-I and IGFBP-3 concentrations in patients’ serum were determined using ELISA kits 10-5600 Active and 10-6600 Active, respectively, from Diagnostic Systems Laboratories (Oxon, United Kingdom). The IGF-I kit procedure contained an acid-ethanol extraction step to separate IGFs from their binding sites. The assays were validated and done according to the instructions of the manufacturer. The molar ratio of IGF-I to IGFBP-3 was calculated as [0.13 × IGF-I concentration (ng/mL)] / [0.036 × IGFBP-3 concentration (ng/mL)].

DNA was extracted from leukocytes or tissues using the Qiagen (Crawley, United Kingdom) kit, and M1dG adduct levels were measured in triplicate by immunoslot blot using a murine monoclonal anti-M1dG antibody provided by Dr. Lawrence Marnett (Vanderbilt University, Nashville, TN) as described previously (30). Goat and anti-mouse horseradish peroxidase–conjugated secondary antibodies were purchased from Dako Cytomation (Ely, United Kingdom). Discrepancies in the amount of DNA per slot were corrected for by staining the nitrocellulose filter with propidium iodide and doing UV light densitometry. The detection limit for M1dG was −1 adduct/107 nucleotides.

**Results**

**Identification of silibinin and metabolites in blood.** Patients with confirmed colorectal cancer received silibinin formulated as silipide at 360, 720, or 1,440 mg daily for a week before colorectal or hepatic surgery (for removal of liver metastases). This intervention was not associated with any adverse effect of silibinin. Silibinin exists as two trans-diastereoisomers (Fig. 1), furnishing two peaks on HPLC analysis with retention times near 21 minutes. All three doses afforded measurable peaks in the plasma at the retention times of authentic silibinin. Figure 2 shows the chromatogram of peripheral plasma from a patient who received the highest dose. These peaks were unambiguously identified as silibinin by MS, as they afforded m/z 481 [M-H+].

**Table 1.** HPLC-MS analysis of species related to silibinin in plasma from humans who had received silibinin (1,440 mg) daily for 7 days as silipide capsules

<table>
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<tr>
<th>m/z [M-H+−]−</th>
<th>Inference</th>
<th>Retention time (min)</th>
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<tr>
<td>481</td>
<td>Silibinin</td>
<td>20.8; 21.4</td>
</tr>
<tr>
<td>657</td>
<td>Silibinin glucuronide</td>
<td>12.6; 15.6; 16.4; 17.8</td>
</tr>
<tr>
<td>833</td>
<td>Silibinin diglucuronide</td>
<td>9.2; 10.8</td>
</tr>
<tr>
<td>561</td>
<td>Silibinin sulfate</td>
<td>17; 17.8; 18.8</td>
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could be detected (Table 1). Single ion monitoring yielded multiple peaks, which could be assigned to silibinin monoglucuronide, silibinin diglucuronide, silibinin monosulfate, and silibinin glucuronide sulfate (Fig. 3). There was also some evidence for the presence of silibinin triglucuronide (m/z 1,008) and O-desmethyl silibinin glucuronide (m/z 643; data not shown). Because of the occurrence of multiple diastereoisomers of silibinin derivatives and given the absence of authentic reference materials, it is impossible to assign specific positional isomers to the multiple peaks seen on single ion monitoring. Figure 2 illustrates that incubation with conjugate-hydrolyzing enzymes furnished a marked increase in the height of the parent silibinin peaks, consistent with the abundant presence of silibinin sulfate and/or glucuronide conjugates.

Silibinin levels in blood and tissues. Extracts of samples of plasma or of normal or malignant colorectal or hepatic tissue were subjected to quantitative HPLC-UV analysis. Peripheral plasma levels were between 0.3 and 4 μmol/L, and they were related to silipide dose (Table 2). Silibinin levels in portal plasma were similar to those in peripheral plasma. Levels of silibinin in normal and malignant colorectal tissue showed considerable variation. They were between 20 and 141 nmol/g and not strictly related to silipide dose. Concentrations of silibinin in hepatic tissue were similar to those in plasma (Table 2).

Effect of silibinin on circulating levels of IGFBP-3, IGF-I, and M1dG. Levels of IGFBP-3, IGF-I, and M1dG were studied as potential markers of silibinin efficacy in the peripheral blood from 24 patients obtained before the first dose of silibinin and between 1 and 4 hours after the last (i.e., seventh) dose. Figure 4 shows that there was no significant difference in concentration of IGFBP-3 or IGF-I between pretreatment and posttreatment serum at any of the dose levels. Neither did comparison of the molar ratio of IGF-I to IGFBP-3 reveal any difference between pre-intervention and post-intervention serum (data not shown). Statistical analysis of the difference between pre-intervention and post-intervention values for IGF-I in serum from patients on 1,440 mg silibinin (Fig. 4B) afforded P = 0.07, tentatively hinting at the possibility that, given a larger cohort of individuals and/or a longer period of intervention, this dose may decrease IGF-I levels. When IGFBP-3/IGF-I levels were compared between patients with different disease stage, there was no obvious difference between Dukes stage on the one side and IGFBP-3/IGF-I levels or susceptibility toward modulation of biomarker levels by silibinin on the other (data not shown).

M1dG levels in leukocytes from peripheral blood isolated before and after the intervention from 20 patients were 3.7 ± 3.7 and 2.3 ± 1.8 adducts per 107 nucleotides, respectively. In the remaining four individuals, leukocytic pre-intervention M1dG levels were below the limit of detection. The values are of the same order of magnitude as those reported previously for human blood (31). Statistical comparison between pre-intervention and post-intervention values of the individual dose groups or the combined doses failed to reveal significant differences, suggesting that consumption of silibinin for a week did not markedly alter leukocytic M1dG. We also compared M1dG levels in normal and malignant tissues obtained by biopsy and after surgery. The values measured in tissue samples were extremely variable between patients, with a substantial number of them close to or at the limit of detection, confounding meaningful interpretation.

Discussion

The outcome of the pilot study described here supports the notion that the repeated administration of silibinin at daily doses up to 1.44 g for a week is safe. This conclusion is consistent with the result of the original evaluation of silipide in human volunteers (12), and a similar inference was made in a preliminary report of a current phase I study of silibinin in hormone-refractory prostate cancer patients, in which up to 20 g silibinin was administered orally daily for a month (32). We describe here for the first time the identification of silibinin plasma metabolites and measurement of silibinin tissue levels in humans who ingested silibinin. Consistent with results obtained previously using liver preparations incubated with silibinin in vitro (27), the results outlined here suggest that silibinin undergoes multiple conjugation reactions in humans. The presence of metabolic conjugates of silibinin in the human biomatrix has hitherto been shown only indirectly (28), in that raised levels of the parent molecule after enzymatic hydrolysis was taken to indicate the presence of conjugates. In contrast, here, the conjugate species silibinin monoglucuronide, silibinin diglucuronide, silibinin monosulfate, and silibinin glucuronide sulfate were unambiguously identified. The silibinin molecule possesses five hydroxy moieties (Fig. 1), three of which are phenolic in nature, but the analysis described here does not allow the exact position of conjugation on the silibinin molecule to be elucidated. On the assumption that the antioxidant activity of silibinin is a function of its polyphenolic structure, the presence of silibinin monosulfate and monoglucuronide, which bear (at least) two intact phenol moieties,

Fig. 3. HPLC-MS selected ion monitoring chromatograms of an extract of a patient’s plasma obtained 3 hours after the last of seven daily doses of silipide (1,440 mg silibinin, administered in three divided portions). The following [M-H+]- ions were monitored: 481 (m/z silibinin; A), 657 (m/z silibinin glucuronide; B), 833 (m/z silibinin diglucuronide; C), 561 (m/z silibinin sulfate; D), and 737 (m/z silibinin glucuronide sulfate; E). For details of dosing and analysis, see Materials and Methods.
suggests that appreciable amounts of circulating silibinin-derived species may share, at least to some extent, the antioxidant potency of the parent molecule.

The plasma levels of silibinin described here need to be compared with those reported previously in healthy volunteers who received oral silipide on a repeated dose schedule (12). In that study, silipide at 720 mg (equivalent to 240 mg silibinin) given daily for 7 consecutive days furnished a mean peak plasma level of 0.38 μmol/L (0.18 μg/mL) reached 0.9 hour after administration of the last dose, and the terminal plasma half-life of the last dose was 3.4 hours. This published data allows a tentative prediction of the levels achieved at the time points at which peripheral blood samples were taken for silibinin analysis in the pilot study described here (between 1 and 4 hours after the last silipide dose). Blood was collected at times that coincide approximately with peak levels on the one side and ~0.9 half-life beyond peak levels on the other. The mean plasma levels for the 360-mg daily doses observed here, which were 0.3 to 0.4 μmol/L (0.14-0.19 μg/mL), are broadly consistent with the earlier volunteer study. The results presented here suggest that the concentration of silibinin achieved after consumption of seven daily doses of up to 1.44 g daily is insufficient to affect circulating levels of IGF-I, IGFBP-3, and M1dG. It is of course conceivable that these putative efficacy biomarkers would be amenable to modulation by these doses of silibinin when given over longer periods of intervention. One week might have been too short to achieve a long-lasting effect on the IGF axis. In athymic mice bearing the DU-145 prostate tumor, administration of silibinin at 0.05% and 0.1% in the diet (equivalent to ~75 and ~150 mg/kg body weight daily, respectively) for the lifetime of animals caused 4- to 6-fold elevation of IGFBP-3 levels over controls, for the two doses, respectively (23). In terms of dose extrapolation based on surface area from mice to humans, these doses are comparable with those used here (see Materials and Methods). Steady-state plasma levels of silibinin, which

<table>
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<tr>
<th>Dose (mg/d)</th>
<th>Peripheral blood levels (μmol/L)</th>
<th>Colorectal tissue levels (nmol/g)</th>
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<tr>
<td></td>
<td></td>
<td>Normal</td>
</tr>
<tr>
<td>360</td>
<td>0.3 ± 0.3*</td>
<td>28 ± 31</td>
</tr>
<tr>
<td>720</td>
<td>0.7 ± 0.6</td>
<td>121 ± 181</td>
</tr>
<tr>
<td>1,440</td>
<td>3 ± 2.3</td>
<td>141 ± 169</td>
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*Mean ± SD of four patients.
accompanied the IGFBP-3-lowering activity in mice, were between 7 and 21 μmol/L (3.4 and 10.1 μg/mL), thus, ~2- to 7-fold above the plasma levels measured in our human pilot study for the 1.44 g (highest) dose of silybin. Intervention with 9-cis-retinoic acid in former smokers was recently reported to cause a significant decrease in serum IGF-1 and an increase in IGFBP-3 (33). It is pertinent to note that the period of intervention in that study was 3 months, substantially longer than the period of intervention with silybin described here. Recently, preliminary evidence for the chemotherapeutic activity of silipide has been published. Repeated daily administration of silipide by oral gavage (at 450 mg/kg silipide, equivalent to 180 mg/kg silybin) caused inhibition of ovarian tumor growth in nude mice, and levels of silybin in the plasma and tumor after termination of the experiment were 15 μmol/L (7.2 μg/mL) and 0.38 nmol/g (0.2 μg/g) tissue, respectively (34). Furthermore, silipide at a dose of 1,800 mg/kg (equivalent to 720 mg/kg silybin) given concomitantly with chemotherapy enhanced the antitumor activity of cis-platinum in a nude mouse model bearing the human A2780 ovarian cancer (35).

Levels of silybin in human colorectal and liver tissue have not been described previously, although silybin-containing remedies have long been marketed as liver protectants. In our study, the colorectal mucosal levels of silybin were highly variable and not related to silipide dose, which may, at least to some extent, be the corollary of the difference between patients in time period (3-6 hours), which elapsed between the consumption of the last dose and surgery. The recommended oral dose of, for example, Legalon (Madaus, Germany), which contains 70 mg silymarin/tablet, is two tablets, taken thrice daily. So the daily dose of silymarin recommended for liver protection is 420 mg. Silymarin contains silydianin and silychristin as well as silybin. On the assumption that silymarin contains ~80% silybin, this silymarin dose would translate into ~340 mg silybin, which is similar to the low dose of silybin administered in the pilot study described here.

Based on this gross calculation, one may tentatively infer that silybin concentrations in liver tissue of an order of magnitude similar to those measured here after the 360-mg dose (i.e., 0.3-0.5 μg/g or 0.6-1 μmol/L in concentration terms) can afford protection of the human liver against toxic insult. In contrast to the relatively low systemic and hepatic levels of silybin, levels achieved in colorectal tissue, between 9.6 and 68 μg/g or 20 to 141 μmol/L in concentration terms, are highly likely to elicit pharmacologic effects in the light of the concentrations reported to cause responses in cells in culture. For example, in cultured DU-145 prostate cancer cells, 15 and 30 μmol/L (7.2-14.5 μg/mL) silybin were sufficient to significantly compromise cell proliferation and increase IGFBP-3 in the cellular supernatant (22).

In summary, repeated administration of silypide achieved levels of silybin in the colorectal tract similar to those known to exert pharmacologic activity. Several silybin sulfates and glucuronides have been identified in human blood, some of them retaining the intact phenol structure, a pharmacophoric feature, which may mediate, at least in part, pharmacologic activity. Intervention for periods of a week seemed to be insufficient for orally consumed silybin to affect the IGF-1/IGFBP-3 system in humans, and circulating silybin-derived species were not abundant enough to reduce blood levels of M$_4$DG significantly. Nevertheless, in the light of the colorectal cancer chemopreventive activity of silybin in rodents (8–10), the high silybin levels achieved in the human colorectal mucosa after consumption of safe silybin doses support its further exploration as a human colorectal cancer chemopreventive agent.

Acknowledgments

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