Chemoprotective Glucosinolates and Isothiocyanates of Broccoli Sprouts: Metabolism and Excretion in Humans

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Abstract

Broccoli sprouts are a rich source of glucosinolates and isothiocyanates that induce phase 2 detoxication enzymes, boost antioxidant status, and protect animals against chemically induced cancer. Glucosinolates are hydrolyzed by myrosinase (an enzyme found in plants and bowel microflora) to form isothiocyanates. In vivo, isothiocyanates are conjugated with glutathione and then sequentially metabolized to mercapturic acids. These metabolites are collectively designated dithiocarbamates. We studied the disposition of broccoli sprout glucosinolates and isothiocyanates in healthy volunteers. Broccoli sprouts were grown, processed, and analyzed for (a) inducer potency; (b) glucosinolate and isothiocyanate concentrations; (c) glucosinolate profiles; and (d) myrosinase activity. Dosing preparations included uncooked fresh sprouts (with active myrosinase) as well as homogenates of boiled sprouts that were devoid of myrosinase activity and contained either glucosinolates only or isothiocyanates only. In a crossover study, urinary dithiocarbamate excretion increased sharply after administration of broccoli sprout glucosinolates or isothiocyanates. Cumulative excretion of dithiocarbamates following 111-μmol doses of isothiocyanates was greater than that after glucosinolates (88.9 ± 5.5 and 13.1 ± 1.9 μmol, respectively; P < 0.0003). In subjects fed four repeated 50-μmol doses of isothiocyanates, the intra- and intersubject variation in dithiocarbamate excretion was very small (coefficient of variation, 9%), and after escalating doses, excretion was linear over a 25- to 200-μmol dose range. Dithiocarbamate excretion was higher when intact sprouts were chewed thoroughly rather than swallowed whole (42.4 ± 7.5 and 28.8 ± 2.6 μmol; P = 0.049). These studies indicate that isothiocyanates are about six times more bioavailable than glucosinolates, which must first be hydrolyzed. Thorough chewing of fresh sprouts exposes the glucosinolates to plant myrosinase and significantly increases dithiocarbamate excretion. These findings will assist in the design of dosing regimens for clinical studies of broccoli sprout efficacy.

Introduction

Numerous epidemiological studies indicate that consumption of large quantities of fruits and vegetables, particularly cruciferous vegetables (e.g., broccoli, cabbage, kale, and Brussels sprouts), is associated with a reduced incidence of cancer (1–6). In a series of laboratory studies, we found that broccoli is rich in phytochemicals that induce phase 2 detoxication enzymes and bolster antioxidant activities in mammalian cells (7–10). Nearly all of this inducer activity is attributable to sulforaphane, which is an isothiocyanate. In intact plants, sulforaphane and other isothiocyanates are stored as inert precursors, termed glucosinolates. Isothiocyanates are released when glucosinolates undergo hydrolysis by myrosinase (E. C. 3.2.3.1), an enzyme that coexists with glucosinolates in crucifers (Fig. 1A). Young broccoli plants are an especially good source of chemoprotective glucosinolates, with levels 20–50 times those found in mature market-stage plants (9). Dietary broccoli sprouts and their component glucosinolates and isothiocyanates induce phase 2 enzymes and afford protection against chemically induced tumors in rodents (8, 9).

The observation that dietary glucosinolates are not detected in urine suggests they are not absorbed (11). Furthermore, with the exception of an early report (12), all available evidence suggests that mammalian tissues cannot convert glucosinolates to isothiocyanates (reviewed in Ref. 13). However, we and others have shown that dietary glucosinolates are indeed converted to isothiocyanates in animals and humans, and that this conversion is mediated by the myrosinase activity of enteric microflora (11, 14, 15). Once generated, isothiocyanates are absorbed and metabolized by sequential enzymatic reactions, the first of which is conjugation with glutathione (Refs. 16 and 17; Fig. 1B). In this report the glutathione adduct and subsequent metabolites are collectively termed dithiocarbamates.

Isothiocyanates and their dithiocarbamate metabolites react quantitatively with 1,2-benzene-dithiol to form 1,3-benzodithiole-2-thione, a cyclic product that is readily separated by
HPLC and detected by UV spectroscopy (11, 18, 19). This cyclocondensation reaction has provided a versatile and highly reproducible method for quantitating glucosinolates and isothiocyanates in plants, as well as their dithiocarbamate metabolites in urine. In previous clinical studies, we found that urinary dithiocarbamates are a sensitive indicator of crucifer consumption, and that dithiocarbamate excretion falls to undetectable levels in the absence of dietary crucifers and cigarette smoking (11). Collective results from dozens of nonsmoking volunteers maintained on diets devoid of glucosinolates or isothiocyanates indicate there are no endogenous sources of urinary dithiocarbamates.4

In a previous series of clinical studies, we made extensive use of the cyclocondensation assay: (a) to prepare known doses of dietary glucosinolates and isothiocyanates, administered in the form of various cruciferous vegetables or condiments; and (b) to quantitate their urinary metabolites (11). In this report, we focus exclusively on 3-day-old broccoli sprouts, which are an unusually rich source of glucoraphanin, the glucosinolate precursor of sulforaphane (structures are shown in Fig. 1A). The objective of these clinical trials was to obtain information on the metabolism and excretion of broccoli sprout glucosinolates and isothiocyanates, information essential for the rational design of dietary intervention studies.

Materials and Methods

This report comprises four human studies: two inpatient protocols with broccoli sprout homogenates (termed HS37 and HS43) and two outpatient protocols with uncooked, fresh, intact broccoli sprouts (HS40 and HS42). For all protocols, the first day of the study is designated day 0.

Healthy Volunteers

Subjects were recruited by advertising on local bulletin boards and were eligible to join if they were 18 years of age or older and in good health. Subjects selected for inpatient studies were within 25% of ideal weight for height and build (20), did not use any form of tobacco or regularly take any medications, had not taken any antibiotics for 4 weeks immediately preceding the study, and had no significant abnormalities on medical history, physical examination, or laboratory tests (including routine hematology; chemistry; urine analysis; hepatitis A, B, and C serologies; and HIV testing). Written informed consent was obtained before enrollment. For each of the four studies, subjects were enrolled and studied in a single cohort.

General Study Design

The studies were conducted in the inpatient and outpatient facilities of the NIH-sponsored General Clinical Research Center in the Johns Hopkins Hospital after approval by the Joint Committee on Clinical Investigation. For inpatient studies, a diet free of glucosinolates or isothiocyanates was provided by the Research Nutrition Service. Every component of the diet was assayed (described below) and found to be free of detectable glucosinolates, isothiocyanates, or phase 2 enzyme-inducing activity. Subjects were asked not to eat or drink anything not provided with their meals and not to use any systemic or topical medications, including skin lotions. Urine was collected throughout the entire study in 8-h collection intervals (7 a.m. to 3 p.m.; 3–11 p.m.; and 11 p.m. to 7 a.m.; unless indicated otherwise) and stored at 4°C for ≤24 h before analysis. On dosing days, subjects consumed nothing but water after midnight; the sprout preparation was administered with 240 ml water at 7 a.m., and breakfast was withheld until 10 a.m. Outpatient studies were conducted similarly, except that volunteers were asked to abstain from dietary sources of glucosinolates and isothiocyanates and to keep a food diary.
Plant preparations were made individually for each of the four studies. All studies were conducted with 3-day-old broccoli sprouts (Brassica oleracea var. italica; cultivar DeCicco). Seeds untreated with pesticides were obtained from Penn State Seeds, Inc. or from Seeds by Design, surface-sterilized with a 1-min rinse in 70% ethanol, exposed for 15 min to 1.3% sodium hypochlorite containing 0.001% Alconox detergent, then rinsed exhaustively with distilled water. Seeds were sprouted in inclined perforated plastic trays (36 × 42 cm), at 25°C without added nutrients, by watering with four 15-s overhead spray cycles per h and maintaining a 16-h light and 8-h dark photoperiod.

Intact Sprouts. Fresh intact sprouts for direct dosing were grown, harvested as described, and stored in the dark at 4°C for no more than 2 days before feeding. Triple solvent (equal volumes of DMSO, dimethylformamide, and acetonitrile) extracts were prepared immediately after harvest (9, 11), evaporated to dryness, then redissolved in distilled water for cyclocondensation, paired-ion chromatography, or assay of inducer activity (described below). Myrosinase activity was assayed in aqueous homogenates, also described below.

Homogenates. An overall scheme for preparing broccoli sprout homogenates suitable for dosing is depicted in Fig. 2. Sprouts were rapidly and gently harvested from the trays, plunged at once into boiling water (1:3.5, w:w) in a 40-liter Groen Model D40 steam-jacketed kettle, and boiled for 5 min. This process effectively destroys the plant myrosinase activity (described below). Myrosinase activity was assayed in aqueous homogenates, also described below.

Preparation and Analysis of Plant Material

Preparation and Analysis of Control Diet

For the inpatient studies, volunteers were maintained on a control diet prepared by the General Clinical Research Center Research Nutrition staff, in a research kitchen located on the ward. The diet was devoid of any spices, herbs, known preservatives, condiments (e.g., mustard, horseradish, wasabi, soy sauce, ketchup, or mayonnaise), crucifers (e.g., broccoli, kale, Brussels sprouts, cabbage, radish, daikon, turnip, kohlrabi, or rutabaga), or vegetables in the onion family (e.g., onions, leeks, garlic, or chives). Salt and sugar were permitted. Breads were prepared from flour free of butylated hydroxyanisole (BHA) or butylated hydroxytoluene (BHT), and meats were baked rather than fried or broiled. Coffee, tea, or caffeinated sodas were not served. Three meals and one snack were served each day, and the menu was repeated every 6 days. At each feeding, the subjects received identical portions, and for HS43, an extra meat or snack was prepared for analytical purposes. Shortly after preparation, the entire meal or snack was homogenized in an equal volume of water and a portion was stored at −20°C until analysis. Beverages and thawed food homogenates were added directly to the assay for quinone reductase-inducer activity (highest concentration of sample, 10% v/v in the assay). Bacterial contamination was successfully avoided by including penicillin and streptomycin in the assays. Limit of detection was <70 units of inducer activity/g of food.5

Analytical Methods

Quinone Reductase Inducer Activity. Induction of quinone reductase catalytic activity was measured in Hepa 1c1c7 murine hepatoma cells grown in microtiter plate wells each containing 150 μl of medium (9, 21, 22). Plant homogenates were centrifuged (5600 × g for 3 min at 25°C), and the supernatant was assayed. The highest concentrations of plant material that doubles quinone reductase specific activity in 48 h (see “Materials and Methods”).

5 The inducer activity of sulforaphane is 33,333 units/μmol; closely related methylthioalkyl isothiocyanates such as isothiocyanate in erinii have about 10% of the potency of sulforaphane; indole isothiocyanates have only about 1% the potency of sulforaphane (7, 25). One unit of inducer activity is defined as the amount of material that doubles quinone reductase specific activity in 48 h (see “Materials and Methods”).
Table 1 Characterization of broccoli sprout dosing preparations

<table>
<thead>
<tr>
<th>Study and dosing preparation</th>
<th>Inducer activity (units/g fresh weight)</th>
<th>Cyclocondensation (μmol/g fresh weight)</th>
<th>Glucosinolates by paired-ion chromatography</th>
<th>Myrosinase (units/g fresh weight)</th>
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<tr>
<td></td>
<td>Total</td>
<td>ITC</td>
<td>Total</td>
<td>ITC</td>
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<tr>
<td>HS37</td>
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<td></td>
<td>Isotiocyanate</td>
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<td>Daikon control</td>
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<td>163,000</td>
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<td>HS40</td>
<td>Intact sprouts</td>
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<td></td>
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<td>9.08</td>
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<td>HS42</td>
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<td>8.43</td>
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<td></td>
<td>Isotiocyanate</td>
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</table>

*ITC, isothiocyanates; GR, glucoraphanin; GI, glucobrassicin; GE, glucorucin; ND, not detectable; -, not determined. See Fig. 3 for glucosinolate structures.

*Myrosinase was added to the assay to determine total activity (glucosinolates plus isothiocyanates); mock reactions without myrosinase were performed to obtain the contribution of isothiocyanates alone.

*Estimation of total glucosinolates excludes minor unidentified peaks, which constitute <5% of the known glucosinolates.

*Glucoraphanin:glucoiberin ratio is 2:1 or more.

homogenate supernatant and urine were equivalent to 0.75 and 15 μl, respectively. Eight or more replicates of eight 1:1 serial dilutions in medium were assayed. To measure total inducer activity in plant samples, excess purified myrosinase (23) and 500 μM ascorbate to activate the myrosinase were added directly to the microtiter plates. One unit of inducer activity is the amount that doubles quinone reductase specific activity in 48 h in a well containing 150 μl of medium.

Cyclocondensation Reaction. This assay detects, with equal efficiency, isothiocyanates and their dithiocarbamate metabolites. Glucosinolates (which do not react in the cyclocondensation reaction) were determined after quantitative conversion to isothiocyanates by treating with purified myrosinase (23) before cyclocondensation. All plant dosing materials and all urine samples were analyzed by cyclocondensation as described in detail previously (11, 18, 19). Just before cyclocondensation, plant homogenates and urine specimens were centrifuged to remove particulate matter (2000 × g for 10 min at 4°C or 200 × g for 5 min at 4°C, respectively). Plant supernatants were maintained for 2 h at 37°C, with or without myrosinase, before cyclocondensation. Because urine samples contain no glucosinolates (11), they were not treated with myrosinase. Cyclocondensation reactions [2 ml; 50% methanol (v/v); 125 mM sodium borate (pH 9.25); 20 mM 1,2-benzene-dithiol; and 5% to 25% (v/v) water] were performed under N₂ in 4-ml glass vials (2 h; 65°C). The vials were loaded into a Waters WISP Autosampler; aliquots of each reaction mixture were injected onto a Whatman Partisil 10 octadecylsilane-2 reverse-phase HPLC column and eluted isocratically with 80% methanol and 20% water (v/v). The area of the 1,3-benzothiolethiole-2-thione peak was integrated at 365 nm by means of a Waters Photodiode Array detector and Waters Millennium software (version 2.15.01). All samples were assayed at three dilutions, and each analytical run included a water blank; a 1,3-benzothiole-2-thione standard curve; and three dilutions of the 3-acetylcysteine derivative of allyl isothiocyanate as a positive control. The lower limits of detection are 80 pmol/g fresh weight broccoli sprouts or 0.2 nmol/ml of urine.

![Paired-ion chromatography of broccoli sprout glucosinolates](image)

Fig 3. Paired-ion chromatography of broccoli sprout glucosinolates. Plant extracts were separated by HPLC in the presence of tetracyclammonium bromide. Glucosinolates (shaded black peaks) were identified by their absence from a repeat chromatogram after treatment of the extract with myrosinase. In order of increasing retention times, the known glucosinolates are glucoraphanin and glucobrassicin (a), 4-hydroxyglucobrassicin (b), glucorucin (c), and neoglucobrassicin (d). Unlabeled minor peaks are tentatively identified as alkyl glucosinolates. Under different experimental conditions, peak a can be resolved into its components; in dozens of such analyses, the glucoraphanin content of broccoli sprouts is reproducibly more than twice that of glucorucin.

Paired-Ion Chromatography of Glucosinolates. Plant extracts were chromatographed in acetonitrile:water (1:1, by volume) containing 5 mM tetracyclammonium bromide on a Whatman Partisil 10 octadecylsilane-2 column (11, 24). This provides a qualitative analysis of the glucosinolate content. To

*J. W. Fahey, unpublished data.
figure 4. time course of dithiocarbamate excretion and quinone reductase-inducer activity in urine of volunteers fed broccoli sprout preparations in a crossover dosing study (hs37). each panel represent the results from a single volunteer. after a 3-day period on control diet, the subjects were fed 111 μmol portions of broccoli sprout isothiocyanates (d) or glucosinolates (g), as indicated; 3 days later, the doses were crossed. on day 9, all subjects were fed 0.92 μmol daikon sprout isothiocyanates (d). urine was collected throughout the entire 12-day study in 4- or 8-h intervals and assayed for dithiocarbamate content (connected ◦) and for quinone reductase inducer activity (bars). the small peak of inducer activity on day 7 for one of the subjects may be an artifact or may reflect a dietary indiscretion.

estimate the quantity of each glucosinolate, an assumption was made that the relative integrated absorbance areas at 235 nm for alkylthioalkylglucosinolates (e.g., glucoraphanin, glucoiberin, and glucoraphin) and indole glucosinolates (e.g., neoglucobrassicin and 4-hydroxyglucobrassicin) are 1.0 and 2.7 times, respectively, those of an equimolar quantity of allyl glucosinolate (9). a representative tracing and glucosinolate structures are presented in fig. 3.

myrosinase activity. uncooked sprouts were homogenized in 33 mM potassium phosphate buffer (pH 7.0) with a Brinkman Polytron homogenizer (0.5× full speed for 15 s at 4°C) and centrifuged (5600 × g for 10 s at 4°C). Myrosinase activity in supernatants was determined by measuring the rate of decrease in absorbance at 235 nm resulting from hydrolysis of allyl glucosinolate, as described previously (23) and with the following modifications. the 1.0-ml reaction mixture contained 33 mM potassium phosphate buffer (pH 7.0), 50 μM allyl glucosinolate, 500 μM ascorbic acid, and 1 mM EDTA. the reaction was initiated by the addition of 50 μl of supernatant. one unit of myrosinase activity hydrolyzes 1 μmol of allyl glucosinolate/min.

statistical methods
results are expressed as means ± SDs. CV is the (SD/mean) × 100%. Student’s t tests are paired and two-tailed.

results

dosing materials and control diet. we made new plant preparations for each of the four studies and analyzed them for (a) inducer activity; (b) glucosinolate and isothiocyanate content by cyclocondensation; (c) glucosinolate profile by paired-ion chromatography; and, in some cases, (d) myrosinase catalytic activity (fig. 2, table 1). as expected, the broccoli sprouts contained high levels of phase 2 enzyme-inducer activity, comparable with those we reported previously (9). by far the greatest contribution to inducer activity is provided by methylthioalkyl glucosinolates, which constitute >93% of the mass of glucosinolates (fig. 3, table 1) and whose cognate isothiocyanates have the greatest inducer potency. 5 indole glucosinolates were present at <7% of total mass (table 1) and afforded only weak inducer potency (25).

Goitrogenic β-hydroxyalkenyl glucosinolates were not detected (fig. 3). multiple independent assays of activity and composition provided consistent results of ~9 μmol glucosinolates/g fresh weight broccoli sprouts (table 1). the inducer potency of broccoli plants depends on numerous factors, including genotype, time of harvest, and conditions of growth (26, 27). the four batches of broccoli sprouts produced over time for these studies were of the same genotype and were grown under controlled environmental conditions, which contributed to the uniformity of the dosing preparations.

During HS43, the control diet was extensively tested for evidence of phase 2 enzyme inducer activity (quinone reductase assay). for all 18 meals and six snacks, inducer activity was 53,000 ± 37,000 units/meal, which is equivalent to <2 μmol of sulforaphane. (by comparison, the glucosinolate/isothiocyanate doses in the four studies reported here ranged from 25 to 200 μmol).

Glucosinolate and Isothiocyanate Crossover Study (HS37). in a 12-day inpatient study conducted with four healthy male volunteers (age range, 43–53 years; 3 blacks and 1 white) we compared the urinary dithiocarbamate excretion after feeding equimolar doses of broccoli sprout glucosinolates or isothiocyanates. Glucosinolate and isothiocyanate dosing preparations (fig. 2) were characterized by bioassay, cyclocondensation reaction, and paired-ion chromatography (table 1). the glucosinolate preparation contained <4% isothiocyanate; conversely, the isothiocyanate preparation contained no detectable glucosinolates. to make the isothiocyanate preparation, we added daikon sprouts (as a source of myrosinase), which themselves contain glucosinolates/isothiocyanates (table 1). however, based on the fact that daikons were added at just 2% by weight, this resulted in a contribution of <1% to the isothiocyanate dosing preparation. after a 3-day washout period, subjects were dosed at 7 a.m. with 111 μmol glucosinolate preparation (two subjects) or isothiocyanate preparation (two subjects). three days later, the doses were crossed; and on day 9 all four subjects were given a volume of daikon homogenate (containing 0.92 μmol isothiocyanates) equivalent to that used to prepare the broccoli sprout isothiocyanate dose. after each dose, urine was collected in 4-h periods, from 7 a.m. to 11 a.m. and 11 a.m. to 3 p.m., and thereafter in 8-h collection periods.
For all four subjects, urinary dithiocarbamate excretion and urinary inducer activity fell below the level of detection during the baseline period (Fig. 4). Thereafter there was a brisk rise in both of these end points in response to dosing with isothiocyanate or glucosinolate preparations. The time course and magnitude of dithiocarbamate excretion closely parallel inducer activity throughout. In all but one case, dithiocarbamate excretion peaked in the 0–4 h postdose collection; for one subject dosed with glucosinolates it peaked in the 4–8 h collection (data not shown). Cumulative 72 h urinary dithiocarbamate excretion after isothiocyanates (corrected for daikon contribution) was significantly greater than that after glucosinolates (88.9 ± 5.5 μmol and 13.1 ± 1.9 μmol, respectively; \( P < 0.0003 \)). Overall, the variation among subjects in excretion after isothiocyanate dosing (CV, 8.2%) was smaller than that after glucosinolate dosing (CV, 17.7%).

Repeated and Escalating Doses of Isothiocyanates (HS43). The reproducibility of dithiocarbamate excretion and the relationship of dithiocarbamate excretion to isothiocyanate dose were evaluated in four male subjects (age range, 21–53 years; two blacks and two whites) who were enrolled in a 20-day inpatient study. A broccoli sprout isothiocyanate preparation was made and characterized as described above (Fig. 2, Table 1), and individual aliquot doses were frozen until use. After a 2-day washout period, subjects were fed 50 μmol of isothiocyanate on days 2, 4, 6, and 8; then 25 μmol on day 11, 100 μmol on day 14, and 200 μmol on day 17. Urine was collected throughout the entire 20-day study, in 8-h intervals.

Three subjects completed the entire study; one withdrew on day 6 just before the third dose. For all subjects, urinary dithiocarbamate excretion fell below detectable levels during the initial control period. After each dose of broccoli sprout isothiocyanates, there was a brisk increase in urinary dithiocarbamate excretion, which peaked (in every case) within the first 8 h after dosing and then decreased in a non-first-order fashion (Fig. 5A). There is remarkably little variation between subjects in the time course of excretion, or in the extent of excretion after (a) repeated doses of 50 μmol (CV ≤ 5.5%; 0–48 h collections); or (b) escalating doses of 25–200 μmol (CV ≤ 9.0%; 0–72 h collections). The intrasubject variability with repeated doses of 50 μmol was similarly low (CV ≤ 7.9%). The dose-response curve is linear over an 8-fold range of doses and extrapolates through the origin (Fig. 5B).

Urinary Dithiocarbamate Excretion after Feeding Intact Sprouts (HS40 and 42). To evaluate the extent to which the hydrolysis of glucosinolates to isothiocyanates may be mediated by the myrosinase activity present in intact uncooked sprouts, in HS40 we monitored urinary dithiocarbamate excretion in a 2-day outpatient study with two male and nine female volunteers (age range, 23–75 years; one Asian, two black, one Hispanic, and seven white). At 7 a.m. on the 2nd day of the study, subjects were fed 11.8 g fresh weight broccoli sprouts (containing 100 μmol of glucosinolates; Table 1) and then given 240 ml of water. Urine was collected continuously for 48 h: three 8-h intervals before dosing and a single 24-h collection after dosing. Two subjects were excluded from additional analysis (one spilled a portion of the 24-h postdosing urine collection; and, based on the food diary, one ate conditions that likely contributed to urinary dithiocarbamates within the 24 h after dosing). Urinary dithiocarbamate excretion fell during the control period from 0.38 ± 0.56 μmol (first baseline) to 0.21 ± 0.34 μmol (third baseline) and rose to 55.8 ± 7.8 μmol during the 24 h after dosing.

To determine whether the extent of conversion of glucosinolates to isothiocyanates is affected by chewing intact uncooked broccoli sprouts, two male and one female subject (age range, 44–75 years; three whites) were enrolled in a 4-day outpatient study (HS42). On the 2nd day, subjects swallowed, without chewing, 12 g fresh weight broccoli sprouts containing 109 μmol of glucosinolates. On the 4th day, they thoroughly chewed before swallowing a 109-μmol dose. Urine was collected continuously throughout the study in three 8-h intervals before each dose and in a single 24-h interval after dosing. Thorough chewing, as compared with swallowing whole sprouts, resulted in significantly greater dithiocarbamate excretion (42.4 ± 7.5 μmol and 28.8 ± 2.6 μmol, respectively; \( P = 0.049 \); Fig. 6).
In previous clinical studies, we evaluated the metabolism and disposition of glucosinolates and isothiocyanates in a variety of crucifers, including market stage broccoli (11). The current studies focus entirely on broccoli sprouts, which differ from mature broccoli plants in two important aspects (9, 13, 25): (a) on a gram-fresh-weight basis, they contain up to 50 times more glucoraphanin, the glucosinolate precursor of sulforaphane, which is the most potent natural phase 2 enzyme-inducer known. Second, they contain substantially less or no detectable amounts of the indole and β-hydroxylkenyl glucosinolates that are associated with potential toxicities (Table 1 and Fig. 3).

The principal end point in these studies was urinary dithiocarbamate excretion, as measured by the cyclocondensation reaction. Although this assay detects with equal efficiency isothiocyanates and each of the sequential metabolites (Fig. 1B), the available evidence indicates that urine contains no isothiocyanates, and that dithiocarbamates are the major metabolites that appear in urine (28–30). Urinary dithiocarbamates therefore reflect absorption, sequential metabolic conversions, and then excretion of the dose. In view of this, the remarkable absence of experimental variability in these studies is particularly striking. We consistently find CVs of ≤10% for dithiocarbamate excretion values that include error contributions from: (a) the cyclocondensation assay; (b) repeated doses in the same individual over time; (c) different individuals; and (d) experiments conducted at different times and with different dosing preparations. This indicates that our methods are highly precise and also suggests that among different individuals, the overall pathway for disposing of dietary isothiocyanates is the same, and that it is operating well below capacity. The latter notion is supported by the strict linear relationship between glucosinolate and phase 2 enzyme-inducer activity (Fig. 4), which affords additional validation of the analytical methods. The inducer activity in these urine samples is quite potent, at ~24,000 units/μmol (as compared with 33,333 units/μmol for sulforaphane; Ref. 7). This activity may be attributable to the dissociation of dithiocarbamates in vitro to yield sulforaphane (31).

Glucosinolates are inert as phase 2 enzyme inducers and must be hydrolyzed to generate the active isothiocyanates. The isothiocyanate and glucosinolate preparations used in this study were designed to represent the upper and lower limits of inducer activity that can be provided by intact broccoli sprouts. The isothiocyanate preparation entirely circumvents the problem of glucosinolate hydrolysis and thus represents a maximum achievable dose of inducer activity, whereas the glucosinolate preparation (which contains no plant myrosinase and depends entirely on gut flora for hydrolysis) represents a minimum achievable dose. We were intrigued to find that total dithiocarbamate excretion after feeding intact uncooked sprouts (~50% of dose) was intermediate between that seen after feeding the limiting glucosinolate or isothiocyanate preparations (~15% or 90% of dose, respectively). This suggests that myrosinase activity in intact sprouts contributes significantly to bioavailability by boosting the glucosinolate-to-isothiocyanate conversion. In view of this finding, it is not surprising that dithiocarbamate excretion was enhanced when fresh sprouts were chewed thoroughly, a process that facilitates hydrolysis by releasing myrosinase and glucosinolates from their sequestered sites within the plant cells.

These findings provide important insights into the disposition of broccoli sprout glucosinolates and isothiocyanates, which will be valuable for the design of rational dosing regimens for clinical studies to evaluate the safety of broccoli sprouts and their efficacy as phase 2 enzyme inducers.

Acknowledgments
We thank the staffs of the Clinical Pharmacology Drug Development Unit and the General Clinical Research Centers for their considerable expertise and help in conducting the clinical work. Jerry Collins and Yuesheng Zhang provided valuable advice.

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