Contribution of gut bacteria to the metabolism of cyanidin 3-glucoside in human microbiota-associated rats

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Abstract

Cyanidin 3-glucoside (C3G) is one of the major dietary anthocyanins implicated in the prevention of chronic diseases. To evaluate the impact of human intestinal bacteria on the fate of C3G in the host, we studied the metabolism of C3G in human microbiota-associated (HMA) rats in comparison with germ-free (GF) rats. Urine and faeces of the rats were analysed for C3G and its metabolites within 48 h after the application of 92 μmol C3G/kg body weight. In addition, we tested the microbial C3G conversion in vitro by incubating C3G with human faecal slurries and selected human gut bacteria. The HMA rats excreted with faeces a three times higher percentage of unconjugated C3G products and a two times higher percentage of conjugated C3G products than the GF rats. These differences were mainly due to the increased excretion of 3,4-dihydroxybenzoic acid, 2,4,6-trihydroxybenzaldehyde and 2,4,6-trihydroxybenzoic acid. Only the urine of HMA rats contained peonidin and 3-hydroxycinnamic acid and the percentage of conjugated C3G products in the urine was decreased compared with the GF rats. Overall, the presence of intestinal microbiota resulted in a 3.7% recovery of the C3G dose in HMA rats compared with 1.7% in GF rats. Human intestinal bacteria rapidly degraded C3G in vitro. Most of the C3G products were also found in the absence of bacteria, but at considerably lower levels. The higher concentrations of phenolic acids observed in the presence of intestinal bacteria may contribute to the proposed beneficial health effects of C3G.

Key words: Cyanidin 3-glucoside: Human microbiota-associated rats: Bacterial conversion: Human intestinal microbiota

Anthocyanins are abundant flavonoids in deep-coloured fruits and vegetables. They have received considerable attention due to their potential health effects. The consumption of anthocyanin-rich foods is associated with a lower risk of all-cause mortality and death due to CHD and CVD. Also, experimental studies have indicated protection by anthocyanins against other age-related disorders, such as cancer and neurodegenerative diseases. The daily anthocyanin intake was estimated to range between 2.7 mg in Germany and 82.5 mg in Finland, with cyanidin glycosides accounting for 44.7% of total intake. Cyanidin 3-glucoside (C3G) is one of the most common anthocyanins contained in dietary plants such as berries. In contrast to other flavonoids, anthocyanins are absorbed as intact glycosides. However, the systemic availability of anthocyanins, their aglycones and the corresponding conjugates is poor. Less than 1% of the ingested anthocyanins are typically absorbed and excreted in the urine. Since the concentration of the parent anthocyanins in the body is low, it has been proposed that their derived metabolites mediate the biological activities and the ensuing health effects of anthocyanins, either directly in the intestine or after absorption. A substantial proportion of ingested anthocyanins enters the large intestine and, thus, gut bacteria affect the fate of anthocyanins. Intestinal microbiota from humans and laboratory animals extensively convert anthocyanins in vitro. Moreover, anthocyanins undergo rearrangements in response to pH. At neutral pH, in particular, anthocyanin aglycones are spontaneously degraded to their corresponding phenolic acids and aldehydes by cleavage of the C-ring. Most studies on the metabolism of C3G in vitro and in vivo have revealed 3,4-dihydroxybenzoic acid (DHBA, protocatechuic acid) as one of the major degradation products. So far, the influence of human intestinal bacteria on the metabolism of C3G and other anthocyanins in vitro is not yet clear.

In the present study, we used rats associated with a human intestinal microbiota (human microbiota-associated (HMA) rats) to investigate the impact of human gut bacteria on the metabolism of C3G. The metabolic fate of C3G in HMA rats

**Abbreviations:** AUC, area under the curve; C3G, cyanidin 3-glucoside; DHBA, 3,4-dihydroxybenzoic acid; GF, germ-free; HCA, 3-hydroxycinnamic acid; HMA, human microbiota-associated; TFA, trifluoroacetic acid; THBA, 2,4,6-trihydroxybenzaldehyde; THBAld, 2,4,6-trihydroxybenzoic acid; UPLC, ultra-performance liquid chromatography; VA, vanillic acid.

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was compared with that of germ-free (GF) rats by analysing urine and faeces. In vitro investigations on the degradation of C3G in the absence and presence of human intestinal bacteria were performed in parallel.

Materials and methods

Chemicals

C3G (purity, 98%) isolated from blackberries by countercurrent chromatography1 was provided by Peter Winterhalter (Institute of Food Chemistry, Technical University of Braunschweig). Cyanidin, peonidin, hippuric acid and phloroglucinol were purchased from Roth. 3-Hydroxyphenylacetic acid, 2-hydroxybenzoic acid, 3-hydroxybenzoic acid, 4-hydroxybenzoic acid, benzoic acid, 3-hydroxycinnamic acid (HCA), ferulic acid, vanillic acid (VA, 4-hydroxy-3-methoxybenzoic acid), 3-methoxybenzoic acid, 2,4,6-trihydroxybenzaldehyde (THBAld), 2,4,6-trihydroxybenzoic acid (THBA), DHBA, 2,4-dihydroxybenzoic acid, 2,6-dihydroxybenzoic acid and catechol were obtained from Sigma-Aldrich. 3-Phenylpropionic acid, 3,4-dihydroxycinnamic acid and phenylactic acid were purchased from Acros. 3-(3-Hydroxyphenyl)propionic acid was isolated from Alfa Aesar.

Degradation experiments with human intestinal bacteria

Faecal suspensions were prepared in an anaerobic workstation (MAKS MG, Meintrup dw scientific) with a gas phase of N₂–CO₂–H₂ (80:10:10, by vol.). Human faeces from one male and three female healthy volunteers, aged 23–45 years and consuming an unspecified Western diet, who had not taken antibiotics for at least 6 months before the study, were each diluted to 1% (w/v) in carbonate–phosphate buffer (0·006 M-carbonate and 0·095 M-phosphate, pH 5·5 (13)).

For incubation experiments with Clostridium saccharogumia DSM 17 460 and Eubacterium ramulus DSM 16 296 (German Collection of Microorganisms and Cell Cultures), the bacteria were grown in a complex medium (standard ST medium (25)). The cells were harvested (8000 g, 3 min, 4°C), washed with carbonate–phosphate buffer and resuspended in the same buffer at concentrations of 1·7 × 10⁸ cells/ml for C. saccharogumia and 1 × 10⁹ cells/ml for E. ramulus.

A 100 μl stock solution of 10 mM C3G in methanol was added to 9·9 ml of bacterial suspensions. Cultures were incubated under a gas phase of N₂–CO₂–H₂ (80:20, v/v) in 16 ml Hungate tubes, fitted with butyl rubber stoppers and screw caps, at 37°C in a water-bath equipped with a magnetic shaker. C3G and bacterial suspensions incubated separately in medium served as controls. Aliquots were taken at different time points, acidified with one volume of 1% aqueous trifluoroacetic acid (TFA, v/v) and immediately lyophilised. At 37°C as described previously (24). The compounds were added from 10 to 100 μM stock solutions in methanol to a final concentration of 100 μM each. Aliquots were taken after 15, 30, 60, 90 and 120 min of incubation, lyophilised and analysed by HPLC.

Rats and treatment

For the present experiment, twenty-nine GF male Sprague–Dawley rats bred in our animal facility from stock obtained from Charles River were used. Experimental conditions were as described previously (24). Whereas seventeen rats were kept under GF conditions, twelve rats were associated at 10 weeks of age with human intestinal microbiota via intragastric application of a 1 ml faecal suspension from a human donor (1·50, w/v, in 0·85% NaCl, w/v) as described previously (24). The composition of the intestinal microbiota in HMA rats was stable throughout the study period (24), which corresponds to data reported in a previous long-term study with rats (25). The microbiological status of GF rats was confirmed throughout the study as described previously (26). The rats were fed a semi-purified flavonoid-free diet during the experiments (24). At 13 weeks of age, 1 ml of 0·1% aqueous citric acid (w/v) was intragastrically administered to each rat as a control. Urine and faeces were collected using metabolism cages (Tecniplast) for 24 h after control application and pooled for every 12 h. C3G at a mean dose of 91·7 (SD 2·3) μmol/kg body weight in 1 ml of 0·1% aqueous citric acid (w/v) was intragastrically applied twice to each rat, at 14 and 20 weeks of age. Following each C3G application, urine and faeces were collected for 48 h, pooled every 12 h. The results obtained for the first and second C3G applications revealed no significant differences. Thus, the two datasets were combined for each rat. At the end of the study, the rats were killed by CO₂ inhalation and pathohistological examinations of liver, lung, kidney, myocardium and spleen were conducted. Furthermore, differential blood counts were determined (Vet-Med-Labor). According to these examinations, weight gain and the overall behaviour, the rats’ health status was not affected by bacterial colonisation or C3G application. The institutional and national guidelines for the care and use of animals were followed and all experimental procedures involving animals were approved by the Office for Agriculture, Ecology and Regional Planning of the State Brandenburg according to § 8.1 Animal Welfare Act.

Preparation of rat samples

Urine was filtered sterile immediately after sampling and acidified with 0.2 volumes of TFA (0·44 M). Until further processing, faeces and acidified urine were stored at –80°C. To extract C3G and its metabolites from the urine, RP-Oasis-HLB cartridges (30 mg, 30 μm; Waters) were used, which had been conditioned by washing with 1 ml methanol and equilibrated by washing with 1 ml of water. After the application of the acidified urine sample (1·2 ml), the cartridge was washed with 1 ml of 1 mM aqueous HCl and 1 ml of 1 mM HCl in 20% methanol. Elution was performed using
Bacterial metabolism of cyanidin 3-glucoside

1 ml of 1 mM-HCl in 90% methanol. The eluates were dried by vacuum centrifugation (RC 10-22, Jouan) and dissolved in the original volume of methanol–water–TFA (90:9:1, by vol.). Faecal samples were thawed, acidified with 0-25 volumes (v/w, relating to wet weight of the original faecal sample) of formic acid (5 M) and lyophilised. Then, eight volumes (v/w) of methanol–HCl (99:1, v/v) were added, and the samples were sonicated for 30 min and centrifuged (7500 g) for 10 min at room temperature. Extraction was repeated five times. The pooled supernatants were dried by vacuum centrifugation. The extraction of urine and faeces was carried out in duplicate. Glucurono- and sulpho-conjugates were determined indirectly by comparative HPLC analysis of samples with and without enzymatic hydrolysis of the conjugated forms using extracts of *Helix pomatia* as described previously (24).

To determine the recovery of C3G, one-fifth of the samples collected during the study were extracted and analysed a second time after spiking with 25 µM C3G. The C3G recovery was 41% for urine and 43% for faeces. In the preliminary experiments, the recovery of cyanidin and peonidin was observed to be comparable with that of C3G. Thus, the values obtained by HPLC analysis of the content of C3G, cyanidin and peonidin in the samples were corrected for the
values obtained by HPLC analysis of the content of C3G, observed to be comparable with that of C3G. Thus, the
experiments, the recovery of cyanidin and peonidin was
was 41 % for urine and 43 % for faeces. In the preliminary
experiments, the recovery of cyanidin and peonidin was observed to be comparable with that of C3G. Thus, the values obtained by HPLC analysis of the content of C3G, cyanidin and peonidin in the samples were corrected for the relative recovery of C3G. DHBA, THBA, VA and THBAld were recovered completely in the preliminary tests.

HPLC analysis The dried samples were each dissolved in the original volume of methanol–water–TFA (90:9:1, by vol.), sonicated at 45 kHz for 5 min (Ultrasonic bath Transonic T 480, Elma) and centrifuged (18 200 g, 5 min). Aliquots (20 µl) of the supernatants were analysed by a Summit HPLC system (Dionex) consisting of a pump (P 680A LPG), an autosampler (ASI-100T), a column oven (TCC-100), a diode-array detector (UVD 340U PDA) and an RP-Zorbax SB-C18 column (5 µm, 4-6 x 150 mm) equipped with a RP-SB-C18 guard column (5 µm) (Agilent Technologies). The column temperature was maintained at 35°C. The mobile phase was a mixture of 0.1 M-aqueous TFA (pH 1, solvent A), methanol (solvent B) and water (solvent C) delivered at 0.1 ml/min with a linear gradient of 0–90% B in 25 min, while solvent C decreased from 90 to 0%. Detection was at 250, 285 and 520 nm. Calibration curves of the corresponding standard compounds were used for quantification of the identified compounds. For unidentified compounds, peak areas were determined at the given wavelength.

Ultra-performance liquid chromatography–electrospray ionisation–MS analysis An ultra-performance liquid chromatography (UPLC) system (Acquity Ultra Performance LC; Waters) consisting of a solvent manager, a sample manager and a diode-array detector was used and connected to a triple quadrupole mass spectrometer with Z-spray atmospheric pressure electrospray ionisation source (Quattro Premier XE; Waters). The column was a UPLC bridged ethyl hybrid (BEH) phenyl column (1-7 µm, 50 mm x 2-1 mm; Waters). The column temperature was maintained at 25°C. The solvents were water–formic acid (95:5, v/v, pH 1-7, solvent A) and methanol (solvent B). They were delivered at 0.35 ml/min with a gradient of 0–10% B in 2-65 min, hold at 10% for 1-46 min and from 10 to 100% in 4-39 min. A 4 µl aliquot of the samples was injected. MS analyses were carried out in positive ionisation mode at a capillary voltage of 0-7 kV, a source block temperature of 110°C and a desolvation temperature of 450°C. The collision gas was Ar at a pressure of 0-30 Pa. The cone voltage was varied between 20 and 60 V and the collision energy between 4 and 35 eV. Data were analysed using MassLynx Software version 4.1 (Waters).

Data evaluation and statistical analysis of the animal study Statistical analysis was carried out using software SPSS 11.5 (SPSS, Inc.). Values were tested for normal distribution using the Shapiro–Wilk test. Time effects were tested by analysing differences between the two C3G applications in GF and HMA rats using a paired t test or, when data were non-normally distributed, by the Wilcoxon test. Independent data were tested for homogeneity of variance by the Levene test and differences were checked for significance using an unpaired t test when data were normally distributed. Differences in non-normally distributed data were checked for significance by the Mann–Whitney test. Differences between the GF and HMA rats were considered significant at P≤0.05. Data are presented as medians.

Results Stability of cyanidin 3-glucoside and cyanidin during simulated gastrointestinal passage The stability of C3G and its aglycone cyanidin during the gastrointestinal passage was tested by incubation for 2 h with simulated gastric and intestinal fluids, respectively. The treatment with simulated gastric fluid decreased C3G and cyanidin concentrations to 75.3 (sd 5.0) and 48.2 (sd 9.1)% of its initial value, respectively. DHBA and THBAld were identified as decomposition products of cyanidin. Incubation with simulated intestinal fluid resulted in a C3G and cyanidin recovery of 76.3 (sd 10.5) and 10.9 (sd 15.5)% respectively.

Conversion of cyanidin 3-glucoside by human intestinal bacteria Faecal slurries prepared from each of the four human donors converted C3G within 2 h completely, while the C3G concentration remained widely stable in the absence of microbiota (90.6 (sd 14.7)% after 24-5 h of incubation; Fig. 1(a)). DHBA, THBAld and four unknown products (P1, P2, P3 and P4) were detected during incubation with faecal slurries (Fig. 1(b)). Of the applied C3G (66.0 (sd 4.4) µM), a maximum of 6-46 (sd 0-98) µM was recovered as DHBA and 1-38 (sd 0-83) µM as THBAld after 10 h of incubation. P1, P2, P3 and P4 were detected as from 30 min of incubation with
human faecal slurries. The peak area of P1 was maximal at 3 h (area under the curve (AUC)285 nm, 2.48 (SD 0.90)). The peak areas of P3 plateaued after 3 h (AUC285 nm, 0.16 (SD 0.07)), of P4 after 8 h (AUC285 nm, 0.12 (SD 0.11)) and of P2 after 24.5 h of incubation (AUC250 nm, 1.41 (SD 0.80)). DHBA was also detected during the incubation of C3G (75.5 (SD 12.4) µM) without faecal microbiota but at a much lower concentration (0.88 (SD 0.18) µM) after 10 h of incubation compared with C3G incubation with faecal slurries (Fig. 1(c)). The concentration of DHBA and THBAld at 10 h of incubation was 1.20 (SD 0.13) and 0.93 (SD 0.11) µM, respectively. The maximum peak areas were detected for P1 after 3 h (AUC285 nm, 1.06 (SD 0.64)) and for P2 after 24 h of incubation (AUC250 nm, 0.75 (SD 0.25)). Cyanidin was as stable as C3G in the absence of faecal microbiota under the conditions used (Fig. 1(a)).

The ability of intestinal bacterial species to convert C3G was tested with E. ramulus and C. saccharogumia, which are known to degrade polyphenolic glycosides (27,28). The incubation of C3G for 48 h with E. ramulus was tested with C. saccharogumia, which, subject to the Cambridge Core terms of use, available at https://www.cambridge.org/core/terms.

The results indicate that C3G was degraded rapidly by bacteria via cyanidin to DHBA, THBAld, P1 and P2. These products could possibly also have been formed spontaneously from cyanidin, in particular at neutral pH inside the bacterial cells. In contrast, P3 and P4 were formed only in the presence of faecal microbiota but not by E. ramulus or C. saccharogumia.

Characterisation of the products formed during bacterial cyanidin 3-glucoside conversion

The spectroscopic properties of the products P1 (λmax = 291/319 nm), P2 (λmax = 316 nm), P3 (λmax = 282/317 nm) and P4

![Image](https://www.cambridge.org/core/10007114512003376)

Fig. 1. (a) Incubation of cyanidin 3-glucoside (C3G) with human faecal slurries (♀), and control incubations of C3G (♂) and cyanidin (◇) without an inoculum. Formation of the products during incubations shown under (a): (b) C3G with human faecal slurries, (c) C3G without an inoculum and (d) cyanidin without an inoculum. C3G, cyanidin, 3,4-dihydroxybenzoic acid (♀) and 2,4,6-trihydroxybenzaldehyde (◇) were quantified by HPLC–diode-array detector based on calibration curves of the corresponding standard compounds. Peak areas were determined for P1 (♀), P3 (△) and P4 (◇) at 285 nm and for P2 (♀) at 250 nm (y-axis on the right). Values are means of faecal slurries of the four tested human donors and, for controls, of three repetitions, with standard deviations represented by vertical bars. AU, arbitrary units.

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(λ\textsubscript{max} = 282/317 nm) differed from those of both C3G (λ\textsubscript{max} = 280/518 nm) and cyanidin (λ\textsubscript{max} = 276/530 nm). In the HPLC analysis, P1 (R\textsubscript{t} = 9.9 min) and P2 (R\textsubscript{t} = 11.7 min) eluted earlier than cyanidin (R\textsubscript{t} = 14.5 min). In addition to their similar UV spectra, P3 and P4 had similar retention times of 16.8 and 16.5 min, respectively.

The four products were further characterised by the electrospray ionisation–MS analysis. The protonated molecule of P1 was most probably represented by m/z 305 [M + H]\textsuperscript{+}, which was supported by further signals at m/z 327 [M + Na]\textsuperscript{+} and m/z 345 [M + K]\textsuperscript{+}. The optimised product-ion spectrum of m/z 305 revealed a base peak at m/z 137 consistent with the B-ring-derived dihydroxybenzoyl cation, while other fragment ions showed less than 8% base peak intensity (Fig. 2). Anthocyanins are known to form chalcones at pH > 3.29,30. Based on its maximum UV absorption at 320 nm, which is typical of chalcones30, and its molecular mass of 304, P1 is proposed to represent the cyanidin-derived α-hydroxychalcone.

The MS spectrum of P2 with signals at m/z 301 [M + H]\textsuperscript{+}, m/z 323 [M + Na]\textsuperscript{+} and m/z 339 [M + K]\textsuperscript{+} indicates a molecular mass of 300. Owing to the limited amount of P2, a product-ion spectrum could not be recorded. P2 may result from the oxidation of P1, with the A-ring forming a quinone methide and the B-ring an 1,2-quinone (for ring designation, see Fig. 6).

The MS analysis of P3 and P4 resulted in complex spectra with postulated protonated molecules [M + H]\textsuperscript{+} at m/z 607 and m/z 605, respectively, and [M + Na]\textsuperscript{+} signals at m/z 629 and m/z 627, respectively. Fragmentation of the protonated molecules of P3 and P4 revealed a pattern of signals similar to each other (Fig. 3(a) and (b)). The difference of two mass units between the molecular peaks and individual fragment ions of P3 and P4 (m/z 589 v. 587, m/z 346 v. 344) indicates that the two compounds differ in their redox state. However, other P3 and P4 specific fragments varied by only one mass unit (m/z 482 v. 481, m/z 464 v. 463). Both the retention times and the spectroscopic behaviour support the close structural similarity of P3 and P4. P3 may be the product of either homodimerisation of P1 followed by the oxidation of one of the four aromatic rings or by heterodimerisation of P1 and an oxidised P1 molecule. P4 could subsequently have been formed by further oxidation of P3. A product of spontaneous cyanidin decomposition with a protonated molecule at m/z 605 [M + H]\textsuperscript{+} was previously proposed to be a dimerisation product of two cyanidin units15.

**Metabolism of cyanidin 3-glucoside in germ-free and human microbiota-associated rats**

The GF and HMA rats excreted C3G, cyanidin, DHBA, THBAld and VA with urine after the oral application of C3G. All compounds also occurred in their conjugated forms except for C3G and cyanidin in HMA rats (Fig. 4). DHBA and VA were the main metabolites in both GF and HMA rats; however, the HMA rats excreted lower proportions as free DHBA and conjugated VA than the GF rats (Fig. 4). Peonidin and HCA in their free and conjugated forms occurred exclusively in the urine of HMA rats but not of GF rats.

The GF and HMA rats excreted C3G, cyanidin, DHBA, THBAld and VA in their free and conjugated forms with faeces throughout the observation period of 48 h after C3G application (Fig. 5). The HMA rats excreted a higher proportion of C3G as free DHBA, conjugated THBAld and THBA in its free and conjugated forms with faeces than the GF rats.

The total excretion of C3G and its metabolites within 48 h after C3G application via the urine and faeces was low in both HMA and GF rats, accounting for 3.7 and 1.7%, respectively, of the ingested C3G dose. The time course of excretion

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**Fig. 2.** Product-ion spectrum of the protonated molecule of the product P1 at m/z 305 [M + H]\textsuperscript{+} obtained at a cone voltage of 20 V and a collision energy of 10 eV.

**Fig. 3.** Product-ion spectra of the protonated molecules of (a) the product P3 at m/z 607 [M + H]\textsuperscript{+} and (b) the product P4 at m/z 605 [M + H]\textsuperscript{+} obtained at a cone voltage of 35 V and a collision energy of 25 eV.
of the identical compounds did not differ between HMA and GF rats. The main excretion route was faeces with 87.8% of total C3G excreted in HMA rats and 68.9% in GF rats, with compounds in their free form largely dominating. The HMA rats excreted more free compounds and conjugates with faeces than the GF rats (Fig. 4), while the opposite was true for the excretion of conjugates with urine (Fig. 4).

Several unknown metabolites were detected after the application of C3G in the urine and faeces of HMA rats, which were not present in the samples of GF rats. These compounds could not be assigned to either the metabolites P1 to P4 observed in the course of in vitro incubation of C3G with human faecal slurries or the following authentic standards: ferulic acid; 3,4-dihydroxycinnamic acid; 3-(3-hydroxyphenyl)propionic acid; 3-phenylpropionic acid; 3-hydroxyphenylacetic acid; phenylacetic acid; 3-methoxybenzoic acid; 2,4-dihydroxybenzoic acid; 2,6-dihydroxybenzoic acid; 2-hydroxybenzoic acid; 3-hydroxybenzoic acid; benzoic acid; catechol; hippuric acid.

Discussion

Compared with other flavonoids, the bioavailability of anthocyanins is very low. Maximum plasma levels of total anthocyanins are in the range of 1–100 nm with doses of 7–1618 μmol in human studies(51). Typically less than 1% of the ingested dose are excreted in the urine by laboratory animals(10), while higher recoveries of up to 5% have been reported after the intake of anthocyanin-rich foods by human subjects(32). However, 60–90% of the anthocyanins were demonstrated to disappear from the gastrointestinal tract within 4 h after a meal, and it has been hypothesised that degradation accounts for this disappearance(51). Anthocyanins are in general stable during incubation with simulated gastric and intestinal fluids(53–56). This was also true for C3G in our hands, whereas its aglycone cyanidin was less stable, resulting in the formation of DHBA and THBAld. The in vitro incubation data and results from studies with ileostomists(12) indicate that significant amounts of dietary anthocyanins reach the large intestine where bacterial degradation may occur. In addition, absorbed anthocyanins are also excreted with bile into the colon(57). Since the contribution of intestinal bacteria to the metabolism of anthocyanins in vivo is still not known, we aimed to elucidate the impact of human gut microbiota on the fate of C3G by comparing the metabolism of C3G in HMA and GF rats. Although urine and faeces were analysed for all the known degradation products derived from C3G so far and other potential metabolites, the total recovery of the applied C3G dose within 48 h was only 3.7% for HMA rats. The minor proportion of 0.4% was excreted with urine, which is in the same range as reported for C3G in previous studies with rodents or human subjects(58–60). When radiolabelled C3G was applied to mice, 3.3% of radioactivity appeared in the urine while 44.5% were recovered in the faeces collected for 24 h after C3G ingestion(61). We recovered only 3.3% of the applied C3G dose in the form of C3G and identified products in the faeces of HMA rats within 48 h. A recovery of 72% of the ingested C3G in the form of DHBA in the blood (44%) and faeces (28%) has been reported in one human study following the ingestion of blood orange juice(62). However, it cannot be excluded that DHBA detected in this study was formed from polyphenolic compounds other than C3G contained in the blood orange juice. In the present rat study with pure C3G, only 1.2% of the dose were recovered as DHBA in the urine and faeces of HMA rats.

Fig. 4. Excretion of cyanidin 3-glucoside (C3G) and its metabolites via the urine of germ-free (GF) rats (n 17, □) and human microbiota-associated (HMA) rats (n 12, ■) within 48 h after the application of C3G. Depicted are median (inner line), 25–75 percentile (box), 10–90 percentile (stamp) and outliers (▲). Values were significantly different between GF and HMA rats: *P<0.05, ***P<0.001. DHBA, 3,4-dihydroxybenzoic acid; HCA, 3-hydroxy- cinnamic acid; THBAld, 2,4,6-trihydroxybenzaldehyde; VA, vanillic acid.

Fig. 5. Excretion of cyanidin 3-glucoside (C3G) and its metabolites via the faeces of germ-free (GF) rats (n 17, □) and human microbiota-associated (HMA) rats (n 12, ■) within 48 h after the application of C3G. Depicted are median (inner line), 25–75 percentile (box), 10–90 percentile (stamp) and outliers (▲). Values were significantly different between GF and HMA rats: **P<0.01, ***P<0.001. DHBA, 3,4-dihydroxybenzoic acid; THBAld, 2,4,6-trihydroxybenzaldehyde; THBA, 2,4,6-trihydroxybenzoic acid.
Interestingly, the recovery of C3G was even low in GF rats corresponding to only 1.7% of the applied dose. Thus, the degradation of C3G by the intestinal microbiota is not the main reason for the low recovery observed in the present and previous studies. The excretion kinetics did not indicate an accumulation of C3G and its products in tissues either in GF or HMA rats. Accumulation of compounds was also excluded in a previous mouse study using radiolabelled C3G. In the present study, several other compounds in addition to DHBA were observed after C3G application in both GF and HMA rats, and further products may have escaped detection. However, the increased faecal levels of DHBA, THBAld and THBA in HMA rats compared with GF rats led to an improved recovery of the C3G dose. This finding and the appearance of peonidin and HCA exclusively in the urine of HMA rats indicate an influence of the intestinal microbiota on the metabolism of C3G in the host.

The present in vitro studies have also shown the accelerated formation of DHBA and THBAld from C3G in the presence of human intestinal bacteria. It may be concluded that the bacteria catalysed the deglycosylation of C3G, which subsequently led to the increased formation of DHBA and THBAld by bacterial and/or spontaneous degradation of cyanidin. The slightly increased urinary levels of cyanidin in HMA rats compared with GF rats and the occurrence of peonidin, the product of cyanidin methylation, exclusively in the urine of HMA rats corroborate this assumption. Cyanidin and its conjugates detected in GF rats were most probably formed by spontaneous decomposition since mammalian glycosidases appear not to act on C3G. Thus, the enzymatic deglycosylation of C3G is catalysed by intestinal bacteria including E. ramulus and C. saccharogumia. The resulting cyanidin is further degraded to DHBA, THBAld and other products depending on the conditions. DHBA and THBAld have been detected after in vitro incubation of C3G with faecal suspensions from human subjects, rats and pigs in previous studies. However, they are also formed through spontaneous decomposition of C3G and cyanidin in the absence of bacteria. This finding is also supported by the present in vitro experiments. The amounts of DHBA and THBAld detected even in vitro were much lower than expected from the disappearance of the parent C3G. Moreover, THBAld appeared at lower levels than DHBA in vitro and in vivo, which has been reported previously and is explained by the increased reactivity of aldehydes.

Following the incubation of C3G with intestinal bacteria, we observed several other products in addition to DHBA and THBAld. For instance, one compound (P1) was tentatively identified as cyanidin chalcone and was also formed spontaneously from cyanidin. Moreover, two other products (P3 and P4) were only formed in the presence of faecal microbiota and appeared to be dimers of the cyanidin chalcone. It is well known that anthocyanidins are stable at pH 1–3 in the form of the flavylum cation but at pH > 4, the carbinol pseudobase, the chalcone pseudobase and the quinoidal base are formed, which may dimerise and/or decompose to the corresponding phenolic acids and aldehydes.

The present results suggest that the formation of C3G products in HMA rats resulted from a combination of conversion steps catalysed by bacterial and host enzymes and spontaneous decomposition (Fig. 6). The methylation of cyanidin to peonidin and the oxidation of THBAld to THBA were only observed in rats but not during fermentation with intestinal bacteria, indicating catalysis of these reactions by host enzymes. VA may result from either the decomposition of peonidin or the methylation of DHBA by the host’s catechol-O-methyltransferase. On the other hand, the intestinal microbiota is able to demethylate peonidin and VA. The mechanism of HCA formation in HMA rats remains to be clarified. Since HCA was not observed as a metabolite of intestinal bacteria in vitro, it is probably formed by the combined activities of bacterial and host enzymes.

In conclusion, phenolic acids, such as DHBA and THBA, formed increasingly in the presence of intestinal bacteria may contribute to the proposed health effects of C3G either directly in the gut or after their absorption from the colon, rather than the parent compound or its unstable derivatives.

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**Fig. 6.** Postulated metabolism of cyanidin 3-glucoside in human microbiota-associated rats. The cyanidin chalcone (P1) was only detected in vitro. Glc, glucosyl moiety.
However, the fate of the majority of the ingested C3G has yet to be explored in future studies. Only the complete knowledge on the absorption and metabolism of C3G and other anthocyanins will allow the elucidation of the mechanism of action of these bioactives.

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