Effects of Human Oral Mucosal Tissue, Saliva, and Oral Microflora on Intraoral Metabolism and Bioactivation of Black Raspberry Anthocyanins

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Abstract

Our oral cancer chemoprevention trial data implied that patient-specific differences in local retention and metabolism of freeze-dried components of black raspberries (BRB) affected therapeutic responsiveness. Subsequent studies have confirmed that anthocyanins are key contributors to BRB's chemopreventive effects. Consequently, functional assays, immunoblotting, and immunohistochemical analyses to evaluate levels and distribution of BRB anthocyanin-relevant metabolic enzymes in human oral tissues were conducted. Liquid chromatography/tandem mass spectrometry (LC/MS-MS) analyses of time course saliva samples collected following BRB rinses were conducted to assess local pharmacokinetics and compare the capacities of three different BRB rinse formulations to provide sustained intraoral levels of anthocyanins. Protein profiles showed the presence of key metabolic enzymes in all 15 oral mucosal tissues evaluated, whereas immunohistochemistry confirmed these enzymes were distributed within surface oral epithelia and terminal salivary ducts. β-Glucosidase assays confirmed that whole and microflora-reduced saliva can deglycosylate BRB anthocyanins, enabling generation of the bioactive aglycone, cyanidin. LC/MS-MS analyses showed retention of parent anthocyanins and their functional, stable metabolite, protocatechuic acid, in saliva for up to 4 hours after rinsing. Furthermore, postrinse saliva samples contained glucuronidated anthocyanin conjugates, consistent with intracellular uptake and phase II conversion of BRB anthocyanins into forms amenable to local recycling. Our data show that comparable to the small intestine, the requisite hydrolytic, phase II and efflux transporting enzymes necessary for local enteric recycling are present and functional in human oral mucosa. Notably, interpatient differences in anthocyanin bioactivation and capacities for enteric recycling would impact treatment as retention of bioactivated chemopreventives at the target site would sustain therapeutic effectiveness. Cancer Prev Res; 4(8); 1209–21. ©2011 AACR.

Introduction

Results from our recent phase I/II pilot chemopreventive trial revealed that approximately one third of the trial participants were “high level responders” (1, 2). As such, their premalignant oral lesions responded to topical application of a gel containing 10% w/w freeze-dried black raspberries (BRB; 0.5-gm gel applied 4 times, total 2 gm daily) by improvement in all parameters assessed including histopathology, gene expression, and reduction in loss of heterozygosity indices (1, 2). The study design accommodated for the extensive interpatient heterogeneity in oral dysplastic lesions by inclusion of pretreatment incisional biopsies to establish pretreatment baseline lesional parameters and provide a confirmed histopathologic diagnosis (1, 2). The beneficial therapeutic responsiveness did not correlate with a lower baseline histologic grade as some of the optimal effects were obtained in persons with higher grade, previously recalcitrant-to-treatment dysplastic lesions (1, 2). These findings imply that patient-specific differences in local pharmacokinetics, that is, target tissue absorption, metabolic bioactivation, and local retention of the BRB constituents affected chemopreventive responsiveness. Subsequent BRB gel pharmacokinetic analyses
conducted in volunteers with healthy oral mucosa, which revealed extensive interparticipant differences with regard to gel absorption, distribution, and local BRB compound retention, support this premise (3). Identification of the pharmacokinetic and metabolic parameters that modulate BRB gel effectiveness could help to optimize chemopreventive strategies. For example, if a BRB metabolite is determined to deliver enhanced chemopreventive effects, then subsequent formulations could incorporate the more bioactive metabolite instead of the parent compound. This approach would provide equal therapeutic benefits to those patients with low bioactivating enzyme activities. Furthermore, identification of key therapy-modulating pharmacokinetic parameters such as absorption or tissue penetration could direct future topical agent formulations, for example, addition of penetration-enhancing compounds. Finally, patient-specific pretreatment metabolic profiling, assessed relative to the identified BRB modulating parameters, could also be used to predict clinical outcomes. Supplemental strategies, such as introduction of additional chemopreventive compounds or different delivery systems, could then be employed on patients’ lesions that show poor uptake and fail to bioactivate BRB chemopreventive compounds.

BRB, which served as the active agents in the gel, contain numerous compounds with potential chemopreventive activities, including anthocyanins, ellagic, ferulic, and p-coumeric acids, vitamins (ascorbic acid, α- and β-carotene, and folate), and minerals (4). It is, however, the BRB anthocyanins, that is, cyanidin 3-rutinoside, cyanidin 3-xylosylrutinoside, cyanidin 3-glucoside, and cyanidin 3-sambubioside, which comprise the predominant (3%-5% BRB weight) polyphenolic compounds in BRB and provide a large component of the chemopreventive impact (5). Our laboratories have shown that the anthocyanin-enriched fraction is the BRB component responsible for inhibition of benzo[a]pyrene diol epoxide (ultimate tobacco-associated carcinogen), redox-mediated activation of the pleiotropic transcription activating factors NF-kB and AP-1 (6). Subsequent studies by Wang and colleagues, which established that BRB anthocyanins prevent esophageal tumors in rats, confirmed the in vivo applicability of Hecht and colleagues in vitro bioassay results (7). Because of their obvious importance in BRB chemoprevention, anthocyanin metabolism is the focus of this study.

Deglycosylation of the parent BRB anthocyanin to its respective aglycone enhances chemopreventive impact by removing the bulky sugar groups which could cause steric hindrance, replacing the glycosidic linkage by another antioxidant-scavenging hydroxyl group and facilitating transport independent cell uptake (8). Previous studies by Walle and colleagues showed that human saliva and oral microflora generate aglycones from intact flavonoid glycosides, for example, genistein, with β-glucosidase representing the key responsible enzyme (9). Additional studies by Fleschut and colleagues showed that gastrointestinal (GI) flora not only metabolize monoglycosylated anthocyanins (such as the recognized less stable cyanidin 3-glucoside) but also deglycosylate the more complex diglycosylated and acetylated anthocyanins (10). In addition to deglycosylation, anthocyanins can be metabolized by phase II enzymes [UDP-glucuronosyltransferases (UGT) and sulfotransferases (SULT)] and also undergo methylation via catechol-O-methyltransferase (COMT; ref. 11). Phase II enzymes are typically associated with enhancing solubility to facilitate compound elimination. Recent investigations by Hu and colleagues, which imply that phase II metabolites of flavonoids undergo enteric recycling within the GI tract, suggest an expanded role for phase II enzymes (12, 13). Oral mucosal anthocyanin recycling would augment chemopreventive efficacy by increasing contact time of the BRB chemopreventives with the target premalignant oral epithelial cells. As the mouth is contiguous with the more distal components of the GI tract, oral cavity enteric recycling is logical and plausible.

Potential contributors to intraoral anthocyanin metabolism include oral tissues, saliva, and oral microflora. Accordingly, this study investigated the effects of these oral components on parameters likely to modulate BRB anthocyanins’ chemopreventive efficacy. Tissue analyses included a comprehensive profiling of enzymes of interest that could affect anthocyanin bioactivation and local intraoral levels. Saliva and oral microflora contributions to anthocyanin metabolism were assessed via pharmacokinetic analyses of 3 different BRB rinse formulations as well as functional activity assays. Collectively, our data support the prospect that enteric recycling of anthocyanins occurs in the mouth. Furthermore, interpatient variations in the extent of this process could directly impact the effects of locally delivered chemopreventives.

Materials and Methods

Participation of human subjects

Human subject participation in these studies was in accordance with Ohio State University Institutional Review Board approval and followed the tenets of the Declaration of Helsinki, 1964. Ten clinically healthy, nonsmoking volunteers between the ages of 19 and 61 participated in the rinse study. These same 10 individuals plus an additional participant provided saliva samples for the functional β-glucosidase activity assay. This 11th saliva donor was also a nonsmoking, clinically healthy male aged 30 years. Fifteen donor tissues, obtained from 15 consented individuals undergoing elective oral maxillofacial surgical procedures were employed for the enzyme metabolic profiling studies by Western blot analyses. As only 10 of the tissues employed for immunoblotting contained adequate tissue for immunohistochemical (IHC) analyses, 5 additional clinically normal oral tissues for the IHC studies were obtained from consented donors. All human subjects participants had uncomplicated medical histories which were characterized by no hospitalizations with the exception of elective procedures, ASA PS1 health status, healthy oral tissues, and no mucosal or intrabony pathologies.
Preparation of the BRB rinse formulations

Three BRB rinse formulations were evaluated in this study (see Table 1 for description of the 3 rinses). All constituents were pharmaceutical grade, and the rinses were prepared fresh on the day of the assay under a laminar flow sterile hood. Rinses were prepared with the same amount of BRB (10% w/w) that was used in our gel formulation (see Table 1; refs. 1–3). Rinse II also included chlorhexidine gluconate (0.12% final concentration). Chlorhexidine gluconate persists in human saliva for at least 12 hours after rinsing (14, 15) and its intraoral sustainability is felt to reflect the cationic salt’s absorption to oral surfaces (14, 15). The purpose for chlorhexidine inclusion was to determine whether or not chlorhexidine could augment BRB absorption and sustainability. Second, as chlorhexidine gluconate is antibacterial, inclusion of chlorhexidine gluconate permitted determination of the relative contribution of salivary and oral tissue metabolizing enzymes in the presence of reduced amounts of oral microflora.

BRB rinse pharmacokinetic analyses

In this cross-over study, 10 participants rinsed with all 3 preparations, with a 7-day “wash out” period between the testing of different rinse formulations (Table 1). Baseline saliva samples (obtained >1 hour following any eating, drinking, or oral hygiene such as tooth brushing or mouth rinses) were obtained before rinsing. Participants vigorously rinsed with 15 mL of the test solution for 3 minutes and volume of the expectorated rinse recorded. Samples were shielded from the light and immediately acidified with formic acid (5% final concentration) followed by storage at −80°C until liquid chromatography/tandem mass spectrometry (LC/MS-MS) analyses. Additional saliva samples were obtained at 15, 30, 60, 120, 180, and 240 minutes after rinsing. All participants refrained from eating, drinking, or oral hygiene during the time course saliva collections. Samples were used to determine the following: (i) prerinse saliva levels of BRB anthocyanins and metabolites, (ii) salivary levels of the parent BRB anthocyanins, the respective aglycone (cyanidin) and the stable phenolic acid end product (protocatechuic acid; PCA) at timed collection points (5, 30, 60, 120, 180, and 240 minutes after rinsing and expectorating with the 3 BRB rinse formulations), (iii) levels of BRB anthocyanins and the aglycone (cyanidin) in the expectorated rinse. Both inter- and intradonor data comparisons of the BRB anthocyanin parent compounds and metabolites were conducted.

Table 1. Composition of BRB oral rinses

<table>
<thead>
<tr>
<th>Purified water BRB (rinse I)</th>
<th>Chlorhexidine gluconate rinse (rinse II)</th>
<th>Augmented rinse + BRB (rinse III)</th>
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<tbody>
<tr>
<td>Purified water</td>
<td>Purified water</td>
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<tr>
<td>BRB (10% final concentration)</td>
<td>Ora-Plus</td>
<td>Ora-Plus</td>
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<tr>
<td></td>
<td>Ora-Sweet SF</td>
<td>Ora-Sweet SF</td>
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<td></td>
<td>Glycerin</td>
<td>Glycerin</td>
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<td></td>
<td>Chlorhexidine gluconate (0.12% final</td>
<td>BRB (10% final concentration)</td>
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<td></td>
<td>BRB (10% final concentration)</td>
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NOTE: Rationale for selection and description of the solution components.
Ora-Plus: This is an oral suspending vehicle, manufactured by Paddock Laboratories. Its composition is as follows: water (97%), sodium phosphate monobasic (<1%), sodium carboxymethylcellulose (<1%), microcrystalline cellulose (<1%), xanthan gum (<1%), carrageenan (<1%). Any other ingredients are less than 0.1% and are considered generally recognized as safe (GRAS). This vehicle is frequently used in pediatric pharmaceutical preparations.
Ora-Sweet SF: This is a sugar and alcohol-free oral syrup vehicle, manufactured by Paddock Laboratories, which is used to provide a sweet fruity odor and taste to liquid medications. Its component ingredients are as follows: sorbitol (10%), glycerin (9%), and sodium saccharin (0.1%). Ora-Sweet SF is used in conjunction with Ora-Plus in the preparation of liquid pediatric pharmaceutical preparations.
Glycerin: This is a GRAS compound that is used as an antimicrobial, preservative emollient. Glycerin was included in the enhanced rinse formulations to augment rinse viscosity and therefore increase opportunity for the adherence of BRB compounds to oral mucosal tissues.
Chlorhexidine gluconate: This is the antimicrobial agent used in oral health care products for treatment of periodontal disease (e.g., Peridex, Gum chlorhexidine gluconate oral rinse). Chlorhexidine gluconate has been shown to show substantivity in the mouth (approximately 30% of chlorhexidine is retained in the oral cavity following rinsing), with the retained drug slowly being released into oral fluids. Substantivity of chlorhexidine gluconate is felt to reflect binding of this positively charged compound to negative charges located on oral mucosal tissues.
Pharmacokinetics of chlorhexidine gluconate: Following rinsing, approximately 30% of chlorhexidine gluconate is retained in the oral cavity, and the retained compound is slowly released into oral fluids. This compound is poorly absorbed from the GI tract, with the mean peak plasma levels of chlorhexidine gluconate obtained at 30 minutes (0.206 μg/g) after ingestion of a 300-mg dose (14).
**Determination of residual levels of the four BRB anthocyanins in postrinse human saliva samples by LC/MS-MS**

The 4 BRB anthocyanins, that is, cyanidin 3-rutinoside, cyanidin 3-xylosylrutinoside, cyanidin 3-sambubioside, and cyanidin 3-glucoside and their aglycone metabolite, cyanidin, in human saliva were quantified in accordance with the method described in our recent article (16). The minimum quantifiable levels for the 4 BRB anthocyanins were 1 ng/mL; the cyanidin level was 2 ng/mL, with standards prepared in 0.2 mL of diluted saliva. The analytic methods were validated for quantification of the BRB anthocyanins and cyanidin in human saliva by using an internal standard (malvidin 3-glucoside) and BRB standards “spiked” with human saliva. Because of the high concentrations of 4 BRB anthocyanins in the early time point samples, these samples were diluted with water to ensure their final concentrations fell within the linear concentration range (1–1,000 ng/mL) of the corresponding standard curve.

**LC/MS-MS analyses to determine the presence of the stable cyanidin degradatory product, PCA, in human saliva following BRB rinses**

Because of the importance of PCA formation in anthocyanin metabolism and bioactivation, a LC/MS-MS method was developed to determine residual PCA levels in post–BRB rinse saliva. Briefly, PCA and the internal standard 4-hydroxy-3-nitrobenzoic acid (HNBA) were obtained from Sigma-Aldrich. The standards in the human samples were analyzed using a API-3000 triple quadrupole mass spectrometer (Applied Biosystems) under an electrospray ionization (ESI) negative mode (standard curves ranged from 1 to 2,000 ng/mL, \( R^2 > 0.99 \)). The minimum quantifiable level for PCA is 1 ng/mL, whereas the detectable level is 0.2 ng/mL in human saliva. Prior to sample analyses, the analytic methods were validated for quantification of PCA in human saliva by using the PCA standard “spiked” with human saliva.

**Determination of β-glucosidase function in human saliva**

Eleven individuals (10 rinse participant donors plus an additional person) provided a baseline (≥3 hours nothing by mouth) saliva sample, then rinsed for 3 minutes with PBS that contained chlorhexidine gluconate (0.12% final concentration). Fifteen minutes after rinsing with the PBS–chlorhexidine solution, another saliva sample was obtained and volumes recorded. Sample β-glucosidase activities were determined by the QuantiChrom β-Glucosidase Assay Kit (BioAssay Systems), which evaluates β-glucosidase hydrolysis of β-nitrophenyl-β-D-glucopyranoside to a chromogenic product. The rate of the reaction is directly proportional to the enzyme activity. β-Glucosidase activity was reported as units per liter (U/L).

**Evaluation of anthocyanin-related metabolic enzymes in human oral mucosa by immunoblot analyses**

Fifteen clinically normal oral mucosal tissues were obtained from consented donors. Tissues were snap frozen in liquid nitrogen and stored at −80°C prior to analyses. Tissues were homogenized in ice-cold radioimmunoprecipitation assay lysis buffer with protease inhibitor cocktail plus phenylmethylsulfonylfluoride (PMSE; Santa Cruz Biotechnology), followed by protein determination (Bio-Rad Bradford protein assay). SDS-PAGE was conducted on 10% acrylamide–bis gels and used the following primary antibodies (Santa Cruz Biotechnology) and working dilutions: mouse anti-β-glucosidase monoclonal antibody (1:200), rabbit anti-COMT polyclonal antibody (1:1,000), rabbit anti-UGT1A polyclonal antibody (1:400), goat anti-UDP-GlcDH polyclonal antibody (1:200), mouse anti-MRP-1 monoclonal antibody (1:100), mouse anti-MRP-2 monoclonal antibody (1:2,000), mouse anti-β-actin monoclonal antibody (1:20,000). Additional antibodies purchased from Abcam were: mouse anti-P-glycoprotein monoclonal antibody (1:2,000), rabbit anti-SGLT1 polyclonal antibody (1:500), rabbit anti-SULT1A1 polyclonal antibody (1:200). Mouse anti-BCRP monoclonal antibody (1:500; Chemicon) was also used. Secondary antibodies were goat anti-mouse IgG-horseradish peroxidase (HRP), goat anti-rabbit IgG-HRP, and donkey anti-goat IgG-HRP, which were purchased from Santa Cruz Biotechnology. Proteins were visualized using ECL Plus Western blotting detection system (GE Healthcare/Amersham) followed by exposure to Kodak films (Kodak) and densitometry analyses (Kodak 1D3 image analysis software; Kodak). Results were normalized relative to endogenous β-actin expression.

**Confirmation of enzyme distribution within the surface epithelium by IHC staining**

Although surface epithelium was present in each specimen, connective tissue stroma was the predominant tissue present in the mucosal specimens. Consequently, tissue-specific IHC analyses were conducted to assess levels of metabolic enzymes at the target tissue epithelial site. Fifteen clinically and histologically normal oral mucosal specimens were used in these studies. The antibodies used for IHC staining were: mouse monoclonal anti-β-glucosidase (1:100) and rabbit polyclonal anti-UGT1A (1:200) antibodies (Santa Cruz Biotechnology), rabbit polyclonal anti-LCT (1:100; Sigma), rabbit polyclonal anti-COMT (1:200; Sigma), mouse monoclonal anti-BCRP (1:20; Abcam), and rabbit polyclonal anti-SGLT1 (1:500; Abcam). Blocking buffer (negative control) was used in place of primary antibody. Samples were incubated with their respective biotinylated secondary antibody (1:200; Vector Laboratories) followed by application of Vectastain ABC reagent (Vector Laboratories). Immunoreactions were visualized using the 3,3’-diaminobenzidine (DAB) substrate, followed by hematoxylin counterstaining. Images were captured using a Nikon DS-Fi-1 high-resolution digital camera and analyzed using Image-Pro Plus 6.2 software (Media Cybernetics). IHC negative controls consisted of the...
exclusive addition of secondary antibody in the absence of primary antibody.

Detection of residual glucuronidated phase II anthocyanin conjugates in postrinse human saliva samples by LC/MS-MS

Studies were conducted to evaluate whether or not anthocyanin conjugates are detectable in saliva following BRB rinses. These initial studies evaluated selected donors’ (based on high levels of salivary anthocyanins) saliva samples from the rinse III group (highest levels of anthocyanin metabolites). Glucuronidation was selected as the initial phase II enzyme pathway to pursue because of the high levels and uniform distribution of UGTs in all of the tissue donors’ mucosa. The LC/MS-MS method employed a Finnigan TSQ Quantum EMR Triple Quadrupole mass spectrometer (Thermo Fisher Scientific Corporation) coupled to a Shimadzu HPLC system (Shimadzu). BRB anthocyanins, cyanidin, and potential glucuronidated conjugates were eluted within 60 minutes using an gradient program with a flow rate 0.20 mL/min on an Aquasil C18 (5 μm) column (2.1 mm i.d. × 150 mm), which consisted of water containing 0.1% formic acid as mobile phase A and acetonitrile containing 0.1% formic acid as mobile phase B. The gradient, which is initially 0% B, was increased to 100% B in 60 minutes and then reverted to 0% B in 1 minute, followed by equilibration at 0% B for 14 minutes, for a total running time of 75 minutes. The tandem mass spectra of the following ions: m/z 625, 757, and 771 corresponding to the molecular ions of glucuronides of C3GLU, C3SAM, and C3RUT, respectively, were utilized.

Determination if saliva and/or human oral microflora generate glucuronidated anthocyanin conjugates in the absence of oral tissues

To evaluate whether or not saliva and/or oral microflora possess functional UGT 1A1 enzymatic activity, ex vivo incubations (37°C, 5% CO2) of 10% BRB with saliva from 6 different donors were conducted. Additional samples which consisted of saliva BRB-supplemented human liver microsomes were also evaluated. These samples were analyzed under the conditions as described in the previous section.

Detection of protocatechuic acid O-glucuronide in post–rinse III saliva samples

As the PCA data showed that rinse III resulted in significantly higher levels of salivary PCA, additional timed harvest post–rinse III saliva samples were collected from 2 donors. The triple-play (Full Scan, Zoom Scan and Collision-induced Dissociation Product ion Scan) LC/MS analysis for the synthetic protocatechuic acid O-glucuronide (PCAOG) standard and samples was conducted using an LCQ system (Thermo) equipped with Shimadzu Class 10Vp HPLC. An Aquasil C18 column coupled to an Aquasil C18 2 mm precolumn filter were used for the separations. The triple-play mode was chosen for analysis of these reconstituted solutions as full mass scan in the range of 150 to 1,000 Th, zoom scan of and data-dependent MS/MS of the most intense peak from the full analytic scan. The mass spectrometer was tuned to its optimum sensitivity by infusion of PCAOG. All operations were controlled by Finnigan Xcaliber software in a Windows NT 4.0 system. For detection of PCAOG in saliva samples, the following transitional ionic reaction parameters established at 329.0 > 153.0 at 35% energy was monitored.

Statistical analyses

The Kruskal–Wallis ANOVA, followed by the Dunns’ multiple comparison test, was used to evaluate the anthocyanin levels at respective time points in the donor population as a whole. Assessment of individual donor responsiveness to the respective rinses, that is, which rinse provided the highest intradonor levels of the parent anthocyanin at time points was determined by the Friedman two-way ANOVA by ranks. A 2-tailed Mann–Whitney U test was used to compare rinse salivary levels of BRB anthocyanins relative to saliva levels obtained from the 10% BRB gel (3). The repeated measures ANOVA, followed by the Bonferroni multiple comparisons test, was used to evaluate salivary retention of PCA. The effects of chlorhexidine gluconate on salivary β-glucosidase activities were assessed via the χ2 1-way classification. The normality of the data distribution determined whether parametric or nonparametric analyses were utilized. Findings with values of P ≤ 0.05 were considered significant.

Results

BRB rinses provide sustained levels of BRB anthocyanins in saliva and show interrinse differences

Postrinse anthocyanin levels showed rinse-related differences, with saliva collected following rinse I containing the highest levels of BRB parent anthocyanins over the time course of the study. Interdonor differences in salivary anthocyanin levels were also apparent. The greatest interdonor variability was noted at the early (0, 5, and 30 minutes) time points, and most apparent in the cyanidin-3- rutinoside (predominant BRB anthocyanin) and PCA (stabilized metabolite) samples. Average interdonor early time point differences were 5-fold, with ranges from 3.6- to 10.3-fold. Interrinse comparisons showed that rinse I provided significantly higher (P < 0.05) BRB anthocyanin levels relative to rinse III at several time points: time 0 (cyanidin-3-rutinoside, cyanidin-3-glucoside, and cyanidin-3-sambubioside), 30 (cyanidin-3-sambubioside), 60 (cyanidin-3-sambubioside), and 240 minutes (cyanidin-3-rutinoside, cyanidin-3-sambubioside). These significant differences were detected in both the inter- and intradonor comparisons. Rinse II anthocyanin levels were only significantly lower than rinse I at 1 time point: time 0, cyanidin-3-glucoside, for both inter- and intra donor comparisons.
of all measured compounds decreased over time, for example, rutinoside levels in saliva ranged from a high (mean ± SD) 2,287.12 ± 489.55 μg/mL at initial time point (rinse I) to 0.05 ± 0.06 μg/mL at the final (240 minute) time point (rinse III). Overall, parent BRB anthocyanins showed the greatest decrease in rinse III over time. In addition, rinse III, which contained the food mimic sweeteners but not the antimicrobial chlorhexidine gluconate, showed the highest levels of anthocyanin metabolism and/or degradation. Finally, while early (up to 5-minute postrinse) saliva samples obtained from all 3 rinse groups showed detectable levels of the aglycone, cyanidin, in all donors, cyanidin levels rapidly decreased and were undetectable in all samples from all rinses by 60 minutes (data not shown).

**BRB rinses provide significantly higher levels of salivary anthocyanins relative to the BRB gel**

Previous studies have confirmed that saliva is the intraoral compartment that retains the highest BRB anthocyanin levels following topical intraoral gel application (3). Gel-to-BRB rinse comparisons of salivary levels of cyanidin 3-rutinoside (most prevalent BRB anthocyanin) and total anthocyanins (collective chemopreventive compounds) were therefore conducted at all shared saliva collection time points (5, 60, 120, and 240 minutes). BRB rinses provided higher salivary levels of both total BRB anthocyanins and the predominant anthocyanin (cyanidin 3-rutinoside) relative to topical gel application (Fig. 1A and B). Notably, 9 of 10 participants retained detectable rutinoside levels at 240 minutes (rinses I and III), and 10 of 10 donors showed detectable rutinoside at 240 minutes after rinse II. In contrast, only 60% (3 of 5) donors retained any salivary anthocyanins, including rutinoside, at the 240-minute time point following gel application.

**The functional antioxidant and stable phenolic acid derivative of BRB anthocyanins (i.e., PCA), persists in the oral cavity following use of BRB rinses**

Sustained intraoral levels of stable BRB metabolites that retain chemopreventive properties could markedly impact chemopreventive efficacy. Pilot studies to assess salivary levels of the antioxidant, stable anthocyanin metabolite, PCA, were therefore also conducted. Our data show that rinse III provided statistically significantly higher levels of PCA over the time course of the assay (P < 0.05, relative to rinse I; P < 0.01, relative to rinse II). At the final (240 minute) time point, 6 of 10 rinse III donors’ saliva contained detectable PCA, whereas only 2 of 10 donors’ saliva from either rinses I or II retained detectable PCA levels (0.2 ng/mL, detection limit).

**Oral microflora, salivary enzymes, and surface oral epithelium can all contribute to intraoral bioactivation of anthocyanins via β-glucosidase activity**

Comparison of saliva samples obtained before and after rinsing with the antimicrobial chlorhexidine gluco-

![Figure 1. Comparisons of the mean BRB anthocyanin levels ± SEM obtained from the 10 participants in the rinse studies relative to data obtained from 5 individuals from the gel pharmacokinetic study (3). A, comparison of salivary levels of total anthocyanins detected over time following rinsing with rinses I, II, or III or topical intraoral placement of a 10% bioadhesive BRB gel. B, comparison of salivary levels of the predominant BRB anthocyanin (cyanidin 3-rutinoside) detected over time following rinsing with rinses I, II, or III or topical intraoral placement of a 10% bioadhesive BRB gel.](cancerpreventionresearch.aacjrns.org)
Finally, high levels of interdonor differences in β-glucosidase activities were observed in both whole saliva samples (including oral microflora, 80-fold) and after rinsing with chlorhexidine gluconate (38-fold).

Results from the Western blot and IHC analyses of oral mucosal tissues confirmed the presence of β-glucosidase protein in human oral mucosa. Western blots also confirmed that all 15 tissues analyzed contained modest to high (18%-66% of control protein) levels of β-glucosidase with a 3.7-fold interdonor protein level differences observed (Table 2). Complementary IHC analyses to identify the sites for cellular distribution revealed β-glucosidase is concentrated in the spinous and granular layers of stratified squamous surface epithelium. Notably, while salivary gland acini and intercalated ducts were negative, terminal salivary ducts showed intense staining for β-glucosidase (Fig. 2). These data imply that the nonmicrobial β-glucosidase reflects contributions of both saliva as well as enzyme release into saliva from both the surface epithelium and the terminal ducts through which saliva passes en route to the surface.

**Oral mucosal tissues possess moderate to high levels of the enzymes that are requisite for flavonoid enteric recycling**

Table 2 depicts a summary of the immunoblotting and IHC profiling of enzymes that can affect anthocyanin metabolism in human oral mucosa. Notably, all tissues evaluated (15 of 15) showed the presence of key phase II drug egress and associated enzymes: UGTs, COMT, UDP-glucose dehydrogenase (UDPGlu-DH), breast cancer resistance protein (BCRP) and β-glucosidase. Fourteen of 15 tissues contained sodium-dependent glucose cotransporter (SGLT1) Interdonor differences in levels of protein expression ranged from 3.5- (UGTs) to 14.2-fold (SGLT1). Concurrent IHC analyses confirmed tissue distributions of 6 of these enzymes (including β-glucosidase) within stratified squamous surface epithelia (Fig. 2) as well as within the terminal ducts of the salivary glands. Although not detected by Western blots, IHC analyses confirmed the presence of lactate phlorizin hydrolase (LPH) in both the stratified squamous surface epithelia and the terminal minor salivary gland ducts. No immunoreactivity was detected in the IHC negative control slides (data not shown). Two additional key enzymes (i.e., arylsulfatase and β-glucuronidase) are known to be present in human saliva (17).

**Glucuronidated anthocyanin conjugates are detectable in post-BRB rinse saliva samples**

To enhance the sensitivity of monitoring for the presence of glucuronidated anthocyanins in these saliva samples, the expected ions of these anticipated glucuronidates at the following m/z values were monitored. As shown in Figure 3A (i), the selected ion chromatograms (SIC) of glucoside (m/z: 625), cyanidin 3-rutinoside (m/z: 771), and cyanidin 3-sambubioside (m/z: 757) in the 5-minute rinse III saliva samples did not reveal any novel peaks that showed the appropriate molecular weight and their product ions corresponding to the glucuronides of cyanidin 3-glucoside and cyanidin 3-rutinoside (m/z: 771; data not shown). There were, however, 3 peaks with the retention time of 19.22, 21.28, and 25.65 minutes labeled as peaks I, II, and III, respectively, in the SIC of cyanidin 3-sambubioside glucuronides detected in 6 of the six 5-minute time point samples and 3 of 6 samples from the 30-minute samples [Fig. 3A (ii)] but absent in the original rinse III solution [Fig. 3A (i)]. Also, the tandem mass spectra of peaks I and II provide the 2 expected product ions of m/z 449 and 287 [Fig. 3A (iii) and (iv)]. These results imply that the components of these 2 peaks are cyanidin 3-sambubioside glucuronides. Further confirmation of these data was obtained following incubation of cyanidin 3-sambubioside with rat liver microsomes, which generated cyanidin 3-sambubioside glucuronides with the similar retention time and tandem mass spectra as previously reported (18). The tandem mass spectrum of peak III showed a single peak of m/z 757.37 [Fig. 3A (v)]. As retention times for glucuronidated anthocyanins are shorter relative to the parent anthocyanins, this peak does not represent cyanidin 3-sambubioside glucuronide. Studies are ongoing to further elucidate the structure of this novel metabolite.

No glucuronidated conjugates of the other BRB anthocyanins were detected within any of the postrinse saliva samples and no cyanidin 3-sambubioside glucuronides were observed after the 30-minute postrinse samples. To test the hypothesis that the detected cyanidin 3-sambubioside glucuronides were generated intracellularly from within oral surface epithelial cells and then transported to saliva, an ex vivo 15-minute incubation of 10% BRB (37°C, 5% CO2) with saliva collected from 6 volunteers was conducted. The mixture was then analyzed under the conditions as described in the section "Detection of residual glucuronidated phase II anthocyanin conjugates in postrinse human saliva samples by LC/MS-MS". Peak I and II were not observed in the saliva incubates of 6 of these 6 samples. Addition of human liver microsomes to the ex vivo saliva samples from these same donors + 10% BRB incubates did result in peaks corresponding to a sambubioside glucuronidated conjugate in 2 of the 6 samples (Fig. 3B).

**PCA glucuronide is also detectable in postrinse saliva**

The total ion chromatogram of synthetic PCAOG (PCAOG generated from the hydrolysis of PCA methyl ester O-glucuronide) in situ showed 1 peak with retention time of 28.12 minutes in the total ion chromatogram (data not shown). The mass spectrum [Fig. 3C (ii)] of PCAOG under negative mode showed 2 predominant peaks with m/z 329.1 and 658.5, corresponding to the deprotonated ion of PCAOG and its dimer, respectively. These data are consistent with a recently reported mass spectrum of PCAOG (19). The data-dependent, collision-induced dissociation spectrum [Fig. 3C (iii)] of the ion of m/z 329.1 has shown the most abundant peak with m/z 153, corresponding to the deprotonated PCA via the labile cleavage of glycosidic bond, along with 2 minor peaks with

---

**Table 2**

<table>
<thead>
<tr>
<th>Enzyme</th>
<th>Interdonor Differences</th>
<th>Data Represented</th>
</tr>
</thead>
<tbody>
<tr>
<td>UGTs</td>
<td>3.5-fold</td>
<td>14.2-fold</td>
</tr>
<tr>
<td>COMT</td>
<td></td>
<td></td>
</tr>
<tr>
<td>UDP-glucose dehydrogenase</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Breast cancer resistance protein (BCRP)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>β-glucosidase</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

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Table 2. Tissue distributions and interdonor variations in oral mucosal anthocyanin-relevant metabolic enzymes

<table>
<thead>
<tr>
<th>Enzyme</th>
<th>Anthocyanin relevant function</th>
<th>Western blot data</th>
<th>IHC data localization</th>
</tr>
</thead>
<tbody>
<tr>
<td>SGLT1</td>
<td>Would facilitate uptake of BRB anthocyanins.</td>
<td>15/15; 14.2-fold (5%–71%)</td>
<td>Surface epithelium, prominent granular layer</td>
</tr>
<tr>
<td>β-Glucosidase</td>
<td>Removes sugar moieties from parent anthocyanins.</td>
<td>15/15; 3.7-fold (18%–66%)</td>
<td>Granular, surface epithelial layer, ducts</td>
</tr>
<tr>
<td>UGTs</td>
<td>Increase water solubility by glucuronide attachment.</td>
<td>15/15; 3.5-fold (66%–228%)</td>
<td>Granular, spinous, epithelial layer, salivary gland, ducts</td>
</tr>
<tr>
<td>Augment local retention via enteric recycling of glucuronidated anthocyanins.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>COMT</td>
<td>Transfers a methyl group to polyphenols, facilitates compound elimination by increasing water solubility.</td>
<td>15/15; 9.3-fold (40%–372%)</td>
<td>Spinous, granular epithelial layer, salivary gland, ducts</td>
</tr>
<tr>
<td>SULTs</td>
<td>Transfers a sulfur moiety, thereby enhancing water solubility.</td>
<td>0/15</td>
<td>N.D.</td>
</tr>
<tr>
<td>UDP-Glu-DH</td>
<td>Generates glucuronides for use by UGTs.</td>
<td>15/15; 7.4-fold (7%–54%)</td>
<td>N.D.</td>
</tr>
<tr>
<td>LPH</td>
<td>Regenerates aglycone from UGT or SULT conjugates.</td>
<td>N.D.</td>
<td>Surface epithelial layer, salivary gland, ducts</td>
</tr>
<tr>
<td>β-Glucuronidase</td>
<td>Removes glucuronides attached by UGTs, functions in enteric recycling.</td>
<td>While N.D. in surface epithelium, β-glucuronidase present in human saliva (17).</td>
<td></td>
</tr>
<tr>
<td>BCRP</td>
<td>ATP-dependent efflux transporter that could participate in the efflux of phase II enzyme conjugates.</td>
<td>15/15; 7-fold (48%–338%)</td>
<td>Spinous, granular, epithelial layer, ducts</td>
</tr>
<tr>
<td>Quinoid anhydrolease</td>
<td>Generates the corresponding phenolic acid and aldehyde from the parent anthocyanin.</td>
<td>N.D.</td>
<td>N.D.</td>
</tr>
<tr>
<td>Permeability glycoprotein (P-gp)</td>
<td>ATP-dependent efflux transporter that could participate in the efflux of glucuronidated compounds.</td>
<td>0/15</td>
<td>N.D.</td>
</tr>
<tr>
<td>MRP1</td>
<td>ATP-dependent efflux transporters that could participate in the efflux of glucuronidated compounds.</td>
<td>15/15 63.4-fold (1.8%–114.2%)</td>
<td>Spinous epithelial layer</td>
</tr>
<tr>
<td>MRP2</td>
<td>ATP-dependent efflux transporters that could participate in the efflux of glucuronidated compounds.</td>
<td>0/15</td>
<td>N.D.</td>
</tr>
</tbody>
</table>

NOTE: Oral mucosal tissue levels and distribution of enzymes associated with anthocyanin metabolism are presented. The Western blot data are organized to depict number of tissues that contain the specific protein, for example, x/15; the fold-difference between the highest and lowest protein levels among the 15 tissues evaluated; and finally, the percentile range in respective protein level expression relative to the housekeeping protein, β-actin. Arylsulfatase is another anthocyanin and local enteric recycling-relevant metabolic enzyme that removes sulfurs attached via SULTs. Because of lack of SULT detection in oral tissues, arylsulfatase-based IHC studies were not conducted. Arylsulfatase removes sulfurs attached via SULTs, functions in local enteric recycling. Because of absence of SULTs in oral tissues, IHC studies not conducted. Arylsulfatase is present in human saliva (17).

Abbreviation: N.D., not detected.

Localization of LPH to surface and terminal duct epithelium and the proportionately low amounts of epithelium relative to connective tissue in oral mucosal biopsies used for the Western blots likely attributes for the discrepancy about negative Western blot data with corresponding positive IHC findings.
m/z 175.0 and 113.2. The ionic transition m/z 329 > 153 was then used to monitor PCAOG to increase the detection limit. Preliminary study showed that this method can detect 100 ng/mL PCAOG with a retention time of 28.12 minutes as shown in Figure 3C (iii). Also, an additional small peak at the retention time of 31.5 minutes was observed, which could be a region isomer of PCAOG. Notably, a peak with the same retention time of 28.2 minutes was detected in the 5-minute saliva sample from 1 of the 2 donors [Fig. 3C (iv)]. Later time point saliva samples (30, 60, 120, and 240 minutes) did not reveal any peaks corresponding to PCAOG in the total ionic chromatogram in saliva samples of either donor [Fig. 3C (v)].

Discussion

Determination of chemopreventive compound levels achieved at the treatment site and correlation between lesional tissue metabolism and therapeutic efficacy are not currently assessed in chemoprevention trials. These aspects, however, are likely among the prime effectors that determine chemopreventive outcomes. The goal of this study was therefore to characterize the anthocyanin-relevant metabolic enzyme profile of oral cavity and to evaluate in situ anthocyanin metabolism.

The entire oral mucosa of persons who develop oral cancer is speculated to have undergone field cancerization (20). Oral cancer chemopreventive strategies should therefore incorporate formulations which address both site-specific and field coverage components. A topical agent plus rinse combination, which would deliver high intrale-sional levels and also disperse chemopreventives throughout the mouth, would fulfill these requirements. Consequently, the purpose of the BRB rinse studies was 2-fold, that is, assess human oral mucosal anthocyanin metabolism and determine whether rinses can provide sustainable chemopreventive levels in the mouth.

Formulation-associated differences in postrinse salivary BRB anthocyanin levels were noted. Rinse I (dH2O only) provided the overall highest postrinse salivary levels of parent BRB anthocyanins. In contrast, rinses II and III, which incorporated flavoring agents, resulted in lower salivary anthocyanin levels. These data suggest that rinses perceived as food stimulated release of saliva and associated salivary β-glucosidase, thereby increasing anthocyanin deglycosylation. Recorded salivary volumes, which showed higher salivary volumes in the rinse II and III samples, support this premise (data not shown). As these data reflect total anthocyanins and account for saliva volumes, the differences do not reflect volume dilutions.
in the rinse II and III samples. Furthermore, the higher anthocyanin levels detected in post-rinse II saliva relative to rinse III, likely reflect chlorhexidine gluconate-mediated reduction of rinse II in oral microflora. The reduced bacterial β-glucosidase and LPH activities in the rinse II samples would decrease deglycosylation, enabling increased retention of parent BRB anthocyanins in saliva.

Ideally, locally delivered chemopreventives persist at the treatment site. Although the saliva data confirmed BRB retention, the real question is whether these levels are clinically relevant. Previous studies have confirmed that the BRB topical gel elicited chemopreventive effects (1, 2) and that the highest levels of the parent BRB anthocyanins were detected in saliva (3). Postrinse BRB salivary anthocyanin levels were therefore compared with salivary levels achieved following gel application. All of the rinses delivered higher salivary BRB anthocyanin levels at the initial time point and also provided greater sustainability over time relative to levels achieved with gel application. The mechanism of rinse-mediated anthocyanin retention is under investigation. For example, chlorhexidine gluconate interacts with salivary films and adsorbs to teeth and oral mucosal surfaces (14, 21). Local adsorption enables retention of approximately 30% of chlorhexidine gluconate, which is then slowly released over time (21). Although BRB anthocyanins are structurally distinct from chlorhexidine gluconate, the postrinse data, which show sustained

Figure 3. A, glucuronidated anthocyanin conjugates are detectable in post-BRB rinse saliva samples. The SIC of the tandem mass spectrometry of the specific ion of m/z 757 in the rinse III solution (i) and the 5-minute saliva sample postrinse with rinse III solution (ii). In contrast to the rinse III solution (i), there are 3 peaks with a retention time of 15, 22, 21, 88, and 25.65 minutes labeled as peaks I, II, and III, respectively, in the SIC of the saliva samples. The tandem mass spectra of peaks I and II provide 2 expected product ions of m/z 287 and 449 (ii and iv), which are consistent with the presence of sambubioside glucuronides. The tandem mass spectrum of peak III showed a single peak of m/z 757.37 (v), which shows a longer retention time relative to sambubioside, represents a novel metabolite. B, the SIC of m/z 757 from the ex vivo incubation (15 minutes) of 10% BRB and human liver microsomes in saliva collected from 1 of 6 healthy volunteers. Similar to data depicted in (A), 3 peaks are also observed in the saliva BRB liver microsomal incubations. There are, however, slight variations in peak retention times and relative intensities which likely reflect different time frames when the samples were analyzed on the column as well as tissue specific variations, that is, liver relative to oral mucosal UGT polymorphisms. C, PCAOG is also detectable in postrinse saliva. The full-scan (i) and collision-induced dissociation (ii) mass spectra of PCAOG under a negative mode showed 2 predominant peaks with m/z 329.1 and 658.8, which correspond to the deprotonated ion of PCAOG and its dimer, respectively. The data-dependent, collision-induced dissociation spectrum of the ion of m/z 329.1 (ii) showed the most abundant peak with m/z 153, corresponding to the deprotonated PCA formed via the labile cleavage of glycosidic bond, and 2 minor peaks with m/z 175.0 and 113.2. The total ion chromatograms (iii) of the ionic transition 329 > 153 of PCAOG (100 ng/mL) in 1% formic acid aqueous solution, the 5-min saliva sample (iv), and the 30-minute saliva sample (v) from a healthy volunteer after rinse with rinse formulation III. A peak with retention time of 28.12 minutes labeled as peak IV and a relative small peak with the retention time of 31.5 minutes labeled at peak V was shown in PCAOG (iii), an identical peak IV was shown in the SIC-5-minute saliva samples but no peaks were shown in the 30-minutes saliva sample (v).
salivary levels of anthocyanins, suggest that BRB anthocya-
nins may also be adsorbed to oral tissues and/or oral
microflora present in plaque. Notably, although the rinses
and gel both contained the identical 10% BRB concentra-
tion, the larger rinse volume delivered a greater amount of
BRB per dose.

Two enzymes, that is, β-glucosidase and LPH are respon-
sible for deglycosylation of anthocyanins, resulting in for-
mation of the aglycone, cyanidin. Our β-glucosidase data,
which showed that oral microflora provide a significant
amount of salivary β-glucosidase activity, are consistent
with previous studies by Walle and colleagues (21).
Furthermore, although the presence of β-glucosidase is
well recognized in human intestinal epithelium (22), only
a single reference that describes β-glucosidase in keratiniz-
ing epithelia was identified (23). Our findings, which show
β-glucosidase protein in human oral mucosa by immuno-
 blotting and confirm the presence of both β-glucosidase and LPH localization in surface oral epithelium by immu-
nohistochemistry, clarify the distribution of these relevant
metabolic enzymes in human oral tissues. These findings
confirm that oral microflora, salivary enzymes, and oral
epithelium can all contribute to generation of the bioactive
glycone from parent anthocyanins.

The strong correlation between smoking and alcohol use
and the development of oral cancer implies that the mouth
is an active site for xenobiotic metabolism. Intraoral xeno-
biotic enzyme characterization studies to date, however,
have focused at the transcriptional level (24). Conse-
quently, little is known about interdonor variations in
protein levels and in which cell population(s) the enzymes
are distributed. Our protein profiles of clinically and his-
tologically normal oral mucosa showed that the majority of
the metabolic and recycling enzymes necessary for antho-
cyanin enteric recycling were present in every oral tissue
specimen evaluated. Furthermore, enzyme distribution, as
determined by IHC analyses, confirmed the relevant
enzymes were almost exclusively distributed within the
statified squamous surface epithelium or within the termi-
 nal salivary ducts (i.e., target tissues). Also apparent were
the fairly extensive interdonor differences in levels of
enzyme present (ranging from 3.5- to 63-fold) and enzyme
activities (over 30-fold differences in β-glucosidase func-
tion). Finally, 2 additional key enzymes essential for recy-
cling of phase II anthocyanin conjugates, that is,
β-glucuronidase and arylsulfatase, are known to be present
in human saliva (17).

Although detection of the necessary bioactivating, phase
II and efflux transporter enzymes in human oral tissues
supports the feasibility for intraoral anthocyanin enteric
recycling, their mere presence does not confirm ongoing
activity. The detection of glucuronidated anthocyanin–
derived conjugates in saliva implies anthocyanin epithelial
uptake, metabolism, and release of a stabilized metabolite
capable of undergoing recycling via deglucuronidation and
cellular reentry (Fig. 4). As UGTs are present in organisms

![Figure 4. Concept of enteric recycling of BRB anthocyanins in oral cavity. SGLT1, which is present in surface oral epithelia, facilitates internalization of the relatively bulky anthocyanins. Intraorally, anthocyanins are deglycosylated via enzymes (β-glucosidase and LPH) which are contained in oral microflora, saliva, and surface oral epithelium. Although the aglycone, cyanidin (depicted in blue dashed line) is a superior antioxidant (due to its additional hydroxyl group), it is unstable and readily degrades to its corresponding phenolic acid, PCA (PCA, shown in red dashed line). PCA, which has a higher chemical and microbial stability relative to cyanidin, is also an antioxidant and may represent the chemical constituent responsible for sustained chemopreventive effects. Although anthocyanins are substrates for SULTs, methylation and glucuronidation reactions represent the primary phase II metabolic routes. Cyanidin and PCA can also undergo glucuronidation. As COMT-mediated methylation occurs at the B ring, cyanidin and PCA should also represent COMT substrates. Both COMT and UGTs are present in human surface oral epithelia.](image-url)
ranging from bacteria to humans, conceptually the detected conjugate could reflect oral bacteria–associated glucuronidation and not entail intracellular uptake, glucuronidation, and egress via UGT efflux transporters (25). Notably, results obtained from the ex vivo saliva BRB incubations, which did not reveal any glucuronidated conjugates in whole saliva without the addition of exogenous liver microsomal enzymes, imply that the requisite UGT1A activity does not exist in saliva. Furthermore, GI microflora–conjugate interactions are commonly associated with deglucuronidation via bacterial β-glucuronidase. Detection of a glucuronidated conjugate is therefore consistent with anthocyanin epithelial uptake (26), metabolism, and release of a metabolite capable of undergoing recycling via deglucuronidation and cellular reentry (Fig. 4). Cyanidin 3-sambubioside, which is found in relatively low levels in BRB (5, 7), was the exclusive glucuronidated anthocyanin conjugate identified. Di- and triglycosylated anthocyanins are more stable than the monoglycosylated forms and are therefore present at higher levels (4, 5). Furthermore, as enteric recycling is a dynamic process, all glucuronidated species are likely present in small quantities and for a very transient time.

Interestingly, our data also confirmed the presence of a PCA glucuronide. These data are consistent with findings by Woodward and colleagues, which imply that PCA glucuronides represent a predominant anthocyanin species following simulated GI digestion (18). Our data imply that the major glucuronidated PCA isomer is formed shortly after rinsing and provide the proof of concept that PCA glucuronides are generated intraorally after anthocyanin administration. Studies to elucidate the intraoral glucuronidation kinetics of intraoral anthocyanin and related compounds are ongoing.

Enterohepatic recycling is a well-accepted mechanism by which parent compounds and their metabolites reenter systemic circulation following first-pass metabolism in the liver. For many tissues and organs such as breast and prostate, this is the exclusive route by which exposure to chemopreventives occurs. In contrast, GI mucosa, which is in direct contact with orally administered agents, avoids first-pass metabolism by the liver and directly absorbs, bioactivates, and metabolizes chemopreventives (27). The requisite components that enable GI enteric recycling of flavonoid compounds are as follows: (i) Hydrolytic enzymes (present in either GI bacteria and/or GI epithelial cells) which deglycosylate parent compounds to generate aglycones, thereby facilitating absorption by the lining epithelial cells. (ii) Intracellular conjugation of aglycones to glucuronic acid or sulfate by intestinal UGTs or SULTs, respectively. (iii) Presence of efflux transporters such as BCRP or multdrug resistance protein (MRP1) which can transport the conjugated metabolites into the intestinal lumen (or oral cavity). (iv) Once in the lumen (or mouth), the conjugates are subject to hydrolysis by salivary and/or bacterial β-glucuronidase or arylsulfatase to regenerate the aglycones or their corresponding phenolic acids and sustain the enteric cycling (12, 28). Our data show that comparable to the small intestine, the requisite hydrolytic, phase II and efflux transporting enzymes are present in the oral cavity.

Although glucuronidation is presumed to be the primary route for anthocyanin phase II metabolism (29), the requisite components for other routes of anthocyanin metabolism, for example, methylation or glutathionylation (COMT and glutathione, respectively) are also present in oral mucosa. Similar to glucuronic acid, conjugation with methyl groups or glutathione could also affect anthocyanin stability and local retention. Notably, interpatient differences in anthocyanin bioactivation and enteric recycling would impact treatment as retention of bioactivated chemopreventives at the target site would sustain therapeutic effectiveness. Pretreatment metabolic profiling of key anthocyanin bioactivating and enteric recycling enzymes and local pharmacokinetics could potentially be used to predict patients most likely to respond favorably to an anthocyanin-based chemopreventive. Such studies are currently ongoing as a component of our BRB gel chemoprevention trial. At the conclusion of the trial, pretreatment profiles will be compared with posttreatment effects to determine the “predictive potential” of anthocyanin metabolic profiles with prospective clinical outcomes.

**Disclosure of Potential Conflicts of Interest**

No potential conflicts of interest were disclosed.

**Acknowledgments**

The authors thank Mary Lloyd and Mary Marin for their expertise in preparation and sectioning of the histologic sections.

**Grant Support**

The work was supported by NIH grants R01 CA129609, RC2 CA148099, and R21 CA132138.

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Received January 24, 2011; revised April 25, 2011; accepted April 27, 2011; published OnlineFirst May 10, 2011.

References


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