Phytochemical-induced mucin accumulation in the gastrointestinal lumen is independent of the microbiota

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Abstract

The mucus layer is critical to gastrointestinal health and ecology. Dietary phytochemicals are well documented to stimulate mucus production and secretion, but the underlying mechanism and effects on gut health are poorly understood. We fed germ-free and conventional mice diets containing approximately 0.4% of polyphenols per gram to determine if the phytochemical-induced accumulation of mucin in the gastrointestinal lumen is dependent on the microbiota. In addition, we assess how increased mucin shapes microbial communities in conventional mice. Germ-free mice receiving a pea (Pisum sativuum) seed coat proanthocyanidin-containing diet (PA) had greater levels of fecal mucin compared to the non-proanthocyanidin-containing (NPA) pea seed coat diet control ($P < 0.05$), confirming that fecal mucin accumulation is independent of the gut microbiota. Conventional mice fed the PA diet and a red osier dogwood (ROD; Cornus sericea) extract diet (DW) had higher mucin levels compared to a control diet without phytochemicals ($P < 0.01$ and $P < 0.05$, respectively). The increase in luminal mucin was associated with consistent increases in bacterial taxa belonging to Lachnospiraceae and [Clostridium] leptum species and a decrease in Romboutsia species. We conclude that phytochemicals have the ability to alter gut microbial ecology by increasing the amount of mucin in the gastrointestinal lumen.
The mucus layer of the mammalian gastrointestinal (GI) tract is a major part of the innate immune system. It is made of heavily glycosylated mucin glycoproteins produced by goblet cells and provides lubrication, hydration, and protection against pathogens and harmful substances that pass through the intestinal environment (1). Two distinctive mucus layers exist: an inner layer devoid of bacteria and an outer loose layer filled with mucolytic and associated microbes that inhabit that particular niche (2–5). Diet has been well recognized to alter host-microbe interactions in the GI tract that affect the integrity of the mucus layer and intestine (6). In particular, phytochemical consumption modulates mucus production, metabolism and has antimicrobial activities that directly affects the gut microbiota; however, the mechanism of their beneficial health outcomes is poorly understood.

Polyphenolic compounds make up the majority of the bioactive phytochemicals consumed in the human diet (7). They are categorized as hydrolysable or condensed (non-hydrolysable) high molecular weight tannins (gallotannins & ellagitannins, and proanthocyanidins, respectively) and low molecular weight polyphenols that include phenolic acids, flavonoids, lignans, stilbenes, and curcumins (8). Because condensed tannins are resistant to host acid and enzyme hydrolysis, they are more likely to reach the gut microbiota compared to the quickly absorbed hydrolysable tannins and low molecular weight phytochemicals. Health benefits have been attributed to their free radical scavenging capacity to neutralize inflammation-causing reactive oxygen species, as well as their direct antimicrobial effect on microbial communities, and the indirect production of bioactive polyphenolic catabolites by the gut microbiota (9,10).
Phytochemicals have an ability to alter mucus physiology, and along with their numerous forms and bioactivities they have shown contradictory effects on resisting pathogen colonization and virulence in the GI tract (11,12). Phytochemical research has focused on improving intestinal barrier integrity with the mechanism of action hinting towards its ability to stimulate the mucus layer, with an increase in mucus production and thickness considered beneficial (11,13,14). However, thickness and expression of mucus-related genes does not necessarily mean that the mucus layer has formed properly for protection. In agreement, a previous study conducted by our group examining the supplementation of peas (Pisum sativuum) rich or low in polyphenol content showed that the polyphenol-rich pea diet increased the amount of fecal mucin in the lumen (12). Excess mucin in the GI lumen was associated with greater Citrobacter rodentium colonization and activation of a proinflammatory response. This diet-induced mucus phenotype has been experimentally tested and confirmed in vitro, as indicated by the ability of galloylated tannins and related compounds to directly cross-link with purified mucins, thereby altering the viscoelastic properties of mucus (15). Therefore, phytochemical consumption increases the accessibility of mucus glycans to the gut microbiota, but to what extent this drives gut ecology has not been determined.

In this study, we investigated whether the presence of microbes is required for the previously observed increase in fecal mucin in response to polyphenol-rich pea seed coat consumption. Whether changes in the gut microbiota drives the mucus phenotype or is a consequence of host-diet interactions remains unknown. We hypothesized that luminal mucin accumulation in the GI tract from phytochemical supplementation is dependent on the microbiota. We fed the proanthocyanidin (PA) and non-proanthocyanidin (NPA) high (20% w/w)-fat pea diets used in our previous study to germ-free (GF) mice and measured fecal mucin
In addition, we tested how the non-hydrolysable PA diet compares to a hydrolysable red-osier dogwood (ROD; *Cornus sericea*) extract on fecal mucin and microbial communities when fed to conventional mice. ROD extracts have been shown in pig models to improve feed efficiency and promote gut resistance to invading pathogens (16); however, the underlying mechanism is poorly understood and possibly driven by changes to the mucus layer. The identification of mucin-degrading bacteria and their impact on the gut niche environment in response to dietary phytochemicals will help determine their contribution to gut ecology and health.

**Material and methods**

**Animals and dietary treatments**

All animal experiments were conducted in accordance with guidelines set by the Canadian Council on Animal Care and approved by the Animal Care and Use Committee at the University of Alberta (Edmonton, AB, Canada). All mice used in this study were bred and maintained in the University of Alberta Axenic Mouse Research Unit. Mice were eight to ten-weeks-old and allowed to acclimatize on an autoclaved standard chow diet (5010 maintenance diet, LabDiet, St. Louis, MO, USA) for a week prior to dietary treatments. Table 1 provides formulations of the isocaloric treatment diets, which were balanced for macronutrients and insoluble fiber with cellulose. Eight female GF Swiss-Webster mice were housed four per open-top cage in the same GF isolator (CEP Standard Safety, McHenry, Illinois, USA) and handling was done directly inside the isolator. GF mice were fed treatment diets containing pea seed coats flours rich (‘Solido’ cultivar; PA) and poor (‘Canstar’ cultivar; NPA) in proanthocyanidin
content as describe previously (12). The diets provided to GF mice were prepared with 15 g instead of 10 g of vitamin mix per kg diet to account for the loss imposed by irradiating the diets to 10 kGY, which was done at the Cross-Cancer Institute at the University of Alberta. Germ free status following the diet intervention was confirmed by anaerobic and aerobic culture of fecal pellets at termination. To investigate the role of microbes and test a second polyphenolic source on the fecal mucin phenotype, we fed a control diet (Control), a PA diet (PA), and the ROD supplemented diet (DW) to conventional Swiss-Webster mice (Table 1). Mice were housed two to four per cage using the Tecniplast Isocage-P bioexclusion system (Buguggiate, VA, Italy) and all animal handling was done in a biosafety cabinet. Control and PA diet groups included two male and two female mice housed separately to determine if sex plays role in the fecal mucin phenotype, previously identified in female mice only (12). The spray-dried ROD extract (Red Dog Enterprises Ltd., Winnipeg, MB, Canada) was added to the dogwood (DW) diet at 4% as done in previous studies (16). ROD extracts can contain bioactive phenolic compounds at 4% to 22% depending on the season (17). The addition of polyphenolic extracts was calculated based on the average total phenolic content of 10% in the ROD extract for the DW diet and 4.51% in the pea seed coat flour for the PA diet. The final amount of proanthocyanidin content in the diets was 0.4% per gram of diet. All diets were prepared aseptically as powdered diet at the University of Alberta and mice had ad libitum access to water and diet throughout the two-week long diet intervention. Body weights were taken every second day and fecal samples were collected aseptically at beginning (day 0) and end (day 14) of the dietary treatment for conventional mice. Mice were euthanized using carbon dioxide. Samples were collected aseptically and stored at -80°C until use.
Table 1. Composition of high (20% w/w)-fat dietary treatments (g/kg).

| Component, g/kg                     | Germ-free |       | Conventional |       |       |
|------------------------------------|-----------|-------|--------------|-------|-------|
|                                    | NPA       | PA    | Control      | PA    | DW    |
| Lard (Tenderflake)                 | 190       | 190   | 190          | 190   | 190   |
| Flaxseed oil                       | 2         | 2     | 2            | 2     | 2     |
| Corn oil (Mazola)                  | 8         | 8     | 8            | 8     | 8     |
| Casein                             | 267       | 262   | 270          | 262   | 267   |
| L-Methionine                       | 2.5       | 2.5   | 2.5          | 2.5   | 2.5   |
| Dextrose                           | 214       | 214   | 214          | 196   | 195   |
| Corn Starch                        | 194       | 194   | 193.4        | 180.7 | 180.4 |
| Cellulose                          | 0         | 0     | 50           | 0     | 45    |
| Mineral Mix                        | 51        | 51    | 51           | 51    | 51    |
| Vitamin Mix                        | 15        | 15    | 10           | 10    | 10    |
| Inositol                           | 6.3       | 6.3   | 6.3          | 6.3   | 6.3   |
| Choline Chloride                   | 2.8       | 2.8   | 2.8          | 2.8   | 2.8   |
| Canstar (NPA) seed coat            | 71.5      |       |              |       | 88.7  |
| Solido (PA) seed coat              |          | 96.5  |              |       |       |
| Red osier dogwood (DW)             |          | 40    |              |       |       |
| Total Weight (g)                   | 1024.1    | 1044.1| 1000         | 1000  | 1000  |
| Fat / g                            | 0.2       | 0.2   | 0.2          | 0.2   | 0.2   |
| Protein / g                        | 0.3       | 0.3   | 0.3          | 0.3   | 0.3   |
| Carbohydrate / g                   | 0.4       | 0.4   | 0.4          | 0.4   | 0.4   |
| Insoluble fiber / g                | 0.04      | 0.05  | 0.05         | 0.05  | 0.05  |
| Energy (At Water) kcal / g         | 4.4       | 4.3   | 4.5          | 4.4   | 4.4   |
| % PACs / g                         | 0         | 0.4   | 0            | 0.4   | 0.4   |

Note: The nutrient content of NPA ('canstar') and PA ('solido') cultivars were analyzed previously and adjusted accordingly. Red osier dogwood extract was provided by Roberts Scales of Red Dog Enterprises Ltd. Diets are isocaloric and contains 0.4% (w/w) of total polyphenols (pea seed coat flour = 4.51%; red osier dogwood extract = 10%).

Fecal mucin assay

Fecal pellets from individual mice were pooled across daily fecal collections throughout the last four days of the two-week dietary treatment. Pooled fecal collections were subsequently
freeze-dried and ground to a powder. A fluorometric assay kit (Fecal Mucin Assay kit; Cosmo Bio co. LTD, Carlsbad, CA, USA) that quantifies N-acetylgalactosamine, the reducing end sugar of the O-linked glycan chain, was used to determine the mucus content (18).

**Microbial community analyses**

Total DNA was extracted from colon contents using the QIamp Fast DNA Stool Mini Kit (Qiagen, Valencia, CA, USA) with an additional bead-beating step using ~200 mg of garnet rock at 6.0 m/s for 60 s on a FastPrep-24 5G instrument (MP Biomedicals, Irvine, CA, USA). Paired-end sequencing was accomplished using the Illumina MiSeq Platform (2 x 300 cycles; Illumina Inc., San Diego, CA, USA). Amplicon libraries were constructed according to the protocol from Illumina (16S Metagenomic Sequencing Library Preparation) that amplified the V3-V4 region of the bacterial 16S rRNA gene: 341F (5’ - TCGTCGGCAGCGTATGCTGTAAGAGACAGCTACGGGNGGCWGGCAG- 3’) and 805R (5’ - GTCTCGTGGGCTCGGAGATGTGTATAAGAGACAGGGTGTATGTAAGAGACAGGACTACHVGGGTATCTAATCC- 3’). Raw sequences were processed with Quantitative Insight into Microbial Ecology 2 (QIIME 2) (19) pipeline using DADA2 to filter, trim and merge paired-end reads into amplicon sequence variants (ASVs). Phylogenetic trees were constructed using the qiime alignment (mafft; mask) and qiime phylogeny (fasttree; midpoint-root) function. Taxonomy was assigned using the qiime feature-classifier classify-sklearn function using the SILVA v138 database trained for the specific amplicon region (20). QIIME2 files (.qza) were imported into R using qiime2R (version 0.99.4) package and analyzed with phyloseq (version 1.34.0) package (21). Sequences belonging to ‘chloroplast’ and ‘mitochondria’ were removed. In addition, ‘Lactococcus’ sequences were
dropped from the analysis because they were suspected to be a contaminant from casein (22).

Numbers assigned to ASVs reflect their total counts from highest to lowest count across samples.

Alpha diversity (Observed, Shannon, phylogenetic diversity (PD)) and beta diversity based on a Bray-Curtis dissimilarity index were done with rarefied reads at a count of 8444.

**Statistical analysis**

Significance testing and graphing for body weights and fecal mucin were done in GraphPad Prism 6 (Graphpad Software, LaJolla, CA. USA) using a t-test or a parametric anova corrected for multiple comparisons with Tukey’s post-hoc test. Data are presented as mean ± standard error of the mean. Letters were used to denote a significance when appropriate.

Statistical significance for alpha diversity was determine with the anova TukeyHSD() correction function. Principal coordinate analyses (PCoA) was plotted using the phyloseq package and clustering significance was determined using the ‘betadisper’ function (23) for dispersion and ‘pairwiseAdonis.dm’ function (24) for orientation. Differential abundance analysis was done with DESeq2 using non-rarefied reads and tree_glom() function to merge similar ASVs. The ‘log2foldchange’ of only the ASVs with a P value less than 0.05 were plotted with bolded ASVs signifying the significant adjusted P value < 0.10, < 0.05 (*), < 0.01 (**) and < 0.001 (***)

**Results**

**Phenolic compounds increase mucin content in the gut**

independently of the microbiota
GF mice fed the PA diet had higher amounts of fecal mucin (P < 0.05) compared to the NPA control diet (Fig 1a). Phytochemicals directly increased fecal mucin in the GI tract independently of the microbiota. The presence of microbes did not alter this outcome as conventional mice displayed a similar increase in fecal mucin in the PA (P < 0.05), as well as the DW (P < 0.05) diet compared to control (Fig 1b). Polyphenolic diets did not affect the weights of conventional mice over the course of the two-week experiment (Fig 1c). Conventional female and male mice fed the Control and PA diets displayed no difference in weight gain or amount of fecal mucin. Although the limited sample size may not adequately represent the sex effect, the impact of phytochemicals on fecal mucin was not different between sexes.
Fig 1. Phytochemical diets (PA and DW) increased the mucin content in the gastrointestinal tract independently of the microbiota. (a) Germ-free Swiss-Webster mice fed the PA diet responded by increasing fecal mucin content, a novel finding that suggest phytochemicals act directly on host mucus chemistry independently of the microbiota (n = 4; * \( P < 0.05 \)). (b) Dietary phytochemicals significantly increased fecal mucin content in PA (\( P < 0.01 \)) and DW (\( P < 0.05 \)) groups of conventional mice (n = 4). (c) Conventional Swiss-Webster mice weights were unaffected by dietary treatment (n = 4).
**Changes in microbial composition in response to phenolic compound rich diets**

Phytochemical diets associated with the fecal mucin phenotype revealed similar changes to microbial communities. PCoA was conducted using Bray-Curtis dissimilarity metric to visualize microbial communities before and after dietary treatment and identify overall differences between treatments. We analyzed microbial communities without separating female and male mice because sex did not appear to alter the amount of mucin recovered in fecal samples. Microbial communities prior to diet intervention were not different between groups (Adonis unadjusted day 0 to control; PA: $R^2 = 0.10$, $P = 0.67$ & DW: $R^2 = 0.11$, $P = 0.56$) but clustered distinctly after 14 days (Adonis unadjusted day 14 to control; PA: $R^2 = 0.27$, $P = 0.11$ & DW: $R^2 = 0.79$, $P = 0.02$). All dietary treatments led to distinct changes in microbial communities when compared to their initial microbial community (Adonis unadjusted day 0 to day 14; Control: $R^2 = 0.53$, $P = 0.05$; PA: $R^2 = 0.55$, $P = 0.04$ & DW: $R^2 = 0.76$, $P = 0.05$) (Fig 2a). Dispersion analysis between groups did not pass significance and shows that the within treatment variability was consistent between groups. PCoA of day 14-microbial communities revealed that both PA and DW diets had similar changes to microbial communities but were still distinct from one another. Principal component (PC) 1, PC2 and PC3 explains 68.2%, 10.3%, and 8.8% respectively and when visualized as PC1 vs PC2 (Fig 2b) and PC2 vs PC3 (Fig 2c) distinct clustering between treatments can be visualized. Alpha diversity metrics of day 0 and day 14 microbial communities show that all dietary treatments reduced the unique counts (Observed; $P < 0.05$); however, diversity as determined by Shannon index revealed that all but the DW group ($P < 0.05$) remained constant compared to both Control and PA groups (Fig 2d).
The PD index revealed that Control and PA diets had lower microbial diversity (P < 0.05) at day 14 compared to day 0, whereas the DW group maintained a similar diversity as at day 0 before diet treatment (Fig 2d).

The phytochemical diets drastically altered the colonic microbiota as determined by differential expression of ASVs using DESeq2 compared to control (Fig 2e-f). This includes numerous ASVs assigned to taxa belonging to the Firmicutes phylum. Compared to the control group, the PA diet reduced *Romboutsia* and *Erysipelatoclostridiaceae*, while increasing *Lachnospiraceae*, *[Clostridium] leptum*, *[Eubacterium] coprostanoligenes*, and a bacterium from the *Clostridia* vadinBB60 group (Fig 2e). The DW diet reduced *Romboutsia*, *Ruminococcaceae* members, *Oscillospiraceae*, *[Eubacterium] coprostanoligenes*, *Peptococcaceae*, and *Ethanoligenenaceae* members of the Firmicute population (Fig 2f). The DW diet increased *Akkermansia muciniphila*, *Parasutterella*, *Alistipes*, *Turicibacter*, and *Bacteroides thetaiotaomicron*, along with an unclassified member from the *Muribaculaceae* family.
Fig 2. Phytochemical diets had a distinct impact on the microbial community structure with consistent changes to Lachnospiraceae, Clostridium, and Romboutsia species in the colon of conventional mice.
Microbial communities prior to diet intervention were not different between groups (Bray-Curtis PCoA; Adonis unadjusted day 0 to control; PA: $R^2 = 0.10$, $P = 0.67$ & DW: $R^2 = 0.11$, $P = 0.56$) but clustered distinctly after 14 days (Bray-Curtis PCoA; Adonis unadjusted day 14 to control; PA: $R^2 = 0.27$, $P = 0.11$ & DW: $R^2 = 0.79$, $P = 0.02$). All mice had similar microbial communities Control: $R^2 = 0.53$, $P = 0.04$; PA: $R^2 = 0.55$, $P = 0.05$ & DW: $R^2 = 0.76$, $P = 0.03$).

At day 14, diet (b) Principal coordinate analysis plot of day 14 microbial communities using principal component 1 (PC1; 68.2%) and principal component 3 (PC3; 8.8%), along with (c) principal component 2 (PC2; 10.3%) plotted against PC3 shows a distinct but subtle similarity between dietary groups. (d) Alpha diversity metrics (Observed, Shannon, PD) of day 0 and day 14 microbial communities shows that all treatment diets reduced diversity ($P < 0.05$); however, DW diet specifically reduced diversity (Observed and Shannon; $P < 0.05$) compared to both Control and PA groups. Differential expression of ASVs as determine by DESeq2 were plotted for (e) PA and (f) DW compared to Control group. Consistent ASVs that respond to PA and DW diets are noted with hashed arrow lines, this includes an increase in Lachnospiraceae and [Clostridium] leptum species along with a decrease in a Romboutsia species ($n = 4$; bolded taxa represent a trend ($P < 0.10$); * $P < 0.05$, ** $P < 0.01$, *** $P < 0.001$).

**Discussion**

Phytochemicals are secondary metabolites produced by plants to communicate with their environment. They are generally considered non-essential nutrients that contribute to physiology through numerous bioactive properties ranging from antioxidant, anti-inflammatory, and antimicrobial to protein chelation and enzyme inhibition (25). Thousands of phytochemicals have been identified and are noted for their free radical scavenging capacity to neutralize reactive
oxygen species associated with inflammation and disease (10). Polymeric phytochemicals, such as proanthocyanidins, are the most abundant and bioactive polyphenols in our diet (26). They are composed of polymeric flavan-3-ol monomers, known as condensed tannins and linked through double A-type and single B-type linkages (26). The degree and type of polymerization, along with the galloylated moieties determine their physicochemical structure and thus their bioactivities on both the gut microbiota and host (27,28). The bioactive properties of polyphenolic compounds and their impact on physiology have made them good candidates for antibiotic alternatives and therapeutics against enteric infection (29).

In this study, we determined that the increase in fecal mucin in response to polyphenol-rich diets observed previously (12,30) occurs independently of the gut microbiota. The inclusion of polyphenols in diet at 0.4% in this study appears safe, as no change in body weight was documented throughout the two-week dietary intervention. This is in accordance with previous studies that show that similar amounts of phytochemicals in diets do not negatively impact growth, and may even improve it (12,16). Despite being quite different with regards to phenolic compound composition, the PA and DW diets both led to increased fecal mucin, suggesting that flavan-3-ol condensed tannins from peas, beans or fruit are not unique in their ability to increase fecal mucin (12,30,31). This occurs independently of the gut microbiota; however, more studies are required to determine the main phytochemicals or groups of chemicals responsible for the mucin phenotype. Our results provide in vivo support to the previous in vitro experiments showing that phytochemicals, particularly the galloylated polyphenols, directly disrupt the viscoelastic properties of mucus by disrupting binding among mucin glycoproteins (15). In our previous study (12), the pea seed coat supplementation (PA diet) led to faster colonization of C. rodentium, a common enteric mouse pathogen, and we suspect that this is a consequence the
direct impact of polyphenols on the mucus layer. Most studies have focused on the beneficial roles that phytochemicals have on health and has been extensively reviewed (32); however, little is mentioned of the phytochemical-mucus interactions in the gut outlined herein. Therefore, in addition to confirming that phytochemical directly increase luminal mucin concentration, an analysis of the colonic microbial community was conducted, with a particular interest in mucolytic microbes that may benefit under these conditions.

Microbial community analysis revealed that pea seed coat and ROD-supplementation alters the gut microbiota. The PCoA plot analyses revealed that the DW diet substantially altered microbial composition compared to the PA diet. Shannon diversity and PD values of the colonic microbiome in the PA group is consistent with our previous experiment (12). The reduced Shannon diversity of the DW group could be explained by the antimicrobial properties of ROD phytochemicals; however, more research is required to confirm the direct antimicrobial actions of ROD supplementation. A study in pigs with a lower inclusion rate of 0.5% ROD extract showed no effect on ileal microbial alpha diversity (Shannon and Simpson) but a prebiotic effect on Lactobacillus species was noted along with no change to growth performance (33). A study in weaned pigs challenged with Escherichia coli k88+ found that 2% and 4% ROD extract diets conferred beneficial effects on growth performance; however, microbial composition was not assessed (16). Phylogenetic diversity in the DW group was maintained at day 14, which could be explained by the increased abundance of Akkermansia muciniphila, Parasutterella and Turicibacter, which only appeared in this DW group at day 14 and were not detected in any group at day 0. Although we did not detect these microbes in the sequencing data of Control and PA groups at day 14, they may have been present below our detection limit for 16s rRNA sequencing. The ROD extract effectively reduced the abundance of some species thereby
opening up a niche for others. For this reason, we see higher PD values in the DW group compared to Control and PA groups.

Microbes that were enriched by both phenolic diets include *Lachnospiraceae* and [*Clostridium* leptum] species, which may reflect a response to increased mucin availability. Luminal mucin levels were confirmed greater in both PA and DW diet groups compared to Control. We characterized an increase in abundance of *A. muciniphila*, a well-known mucolytic microbe, in the DW group but not the PA group. Mucin supplementation has been confirmed in mice to encourage mucin degrading bacteria, such as *A. muciniphila*, and mitigates diet-induced microbiota perturbations (34). Moreover, phytochemicals are well-known to increase the abundance of *A. muciniphila* and improved health outcomes (35). The phytochemical-induced mucin phenotype may partly explain their increased abundance in the gut; however, the absence of *A. muciniphila* in the PA diet suggests other factors contributed to their increase in the DW group. A study in mice using jaboticaba fruit, which is high in flavan-3-ols, at 5%, 10%, and 15% of diet found an increase in *A. muciniphila* at only 10% and 15% (36) suggesting a dose-dependent threshold that supports their growth likely exists in the gut. The lack of *A. muciniphila* in the PA group of this present study is inconsistent with our previous study (12) and suggest mice did not receive the necessary dose of polyphenols to encourage *A. muciniphila* fitness in the gut. The moderate effect of the PA diet on microbial communities could be explained a loss in the antimicrobial actions of pea phytochemicals after long-term storage. As a result, the PA diet had less severe impacts on microbial community structure as compared to the DW diet and suggested that the interactions among microbes are stable enough to prevent *A. muciniphila* expansion even with increased access to mucin glycans. In addition, both PA and DW diets led to a consistent decrease in abundance of *Romboutsia*, a species that is highly adapted to nutrient-
rich environments (37) and a potential marker of stability in the gut. The competitive advantage gained by mucolytic bacteria may have altered the nutrient-rich niches in the gut that genera like *Romboutsia* depend on for growth. The mucus layer supports microbial niches by directly providing glycans for energy and indirectly through cross-feeding from one microbe to another (5). Future research should focus on characterizing the bioactive compounds promoting luminal mucin accumulation, as well as identifying the type and source of the accumulating mucins (1). Further knowledge of these interactions will provide the foundational framework necessary to understand how the mucus layer contributes to host-microbe stability and health.

**Conclusion**

The production and maintenance of the mucus layer is a vital part of intestinal homeostasis. It has become clear that host mucus provides a foundation of host derived glycans that supports mutualism and commensalism among microbes in the gut. Understanding how phytochemicals influence the viscoelasticity of the mucus layer will help to determine the best therapeutic use of phytochemicals to promote health. This research provides insight into establishing the mechanisms involved in the ability of mucus to stabilize gut ecology and control microbial communities. Further studies are required to determine the specific phytochemical compounds and structure that induce mucus secretion and/or disrupt mucin binding and formation.

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