Conversion of rutin, a prevalent dietary flavonol, by the human gut microbiota

Alessandra Riva¹, Ditta Kolimár², Andreas Spittler³, Lukas Wisgrill⁴, Craig W. Herbold¹, László Abránkó², David Berry⁵

¹Centre for Microbiology and Environmental Systems Science, Department of Microbiology and Ecosystem Science, Division of Microbial Ecology, University of Vienna, Vienna, Austria, Austria, ²Faculty of Food Science, Department of Applied Chemistry, Szent István University, Budapest, Hungary, ³Core Facility Flow Cytometry & Department of Surgery, Research Lab, Medical University of Vienna, Vienna, Austria, Austria, ⁴Division of Neonatology, Pediatric Intensive Care and Neuropediatrics, Department of Pediatrics and Adolescent Medicine, Medical University of Vienna, Vienna, Austria, Austria, ⁵Joint Microbiome Facility of the Medical University of Vienna and the University of Vienna, Austria

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The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Author contribution statement

DB, LA and AR conceived and designed the experiments. AR performed the experiments and data analyses. AR and DK performed anaerobic incubation experiments. AS, LW and AR performed FACS sorting. CH performed bioinformatics analyses. AR and DB wrote the paper. All authors have given approval to the final version of the manuscript.

Keywords

dietary bioactives, Rutin, Gut Microbiota, Fluorescence activated cell sorting (FACS), rutin metabolism, inter-individual variability

Abstract

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The gut microbiota plays a pivotal role in the conversion of dietary flavonoids, which can affect their bioavailability and bioactivity and thereby their health-promoting properties. The ability of flavonoids to stimulate the activity of the microbiota has, however, not been systematically evaluated. In the present study, we used a fluorescence-based single-cell activity measure [biorthogonal non-canonical ammino acid-tagging (BONCAT)] combined with fluorescence-activated cell-sorting (FACS) to determine which microorganisms are stimulated by the flavonoid rutin. We performed anaerobic incubations of human fecal microbiota amended with rutin and in the presence of the cellular activity marker L-Azidohomoalanine (AHA) to detect rutin-stimulated cells. We found that 7% of cells in the gut microbiota were active after a 6 h incubation and 23% after 24 h. We then sorted BONCAT-positive cells and observed an enrichment of Lachnospiraceae (Lachnoclostridium, and Eisenbergiella), Enterobacteriaceae, Tannerellaceae and Erysipelotrichaceae species in the rutin-responsive fraction of the microbiota. There was marked inter-individual variability in the appearance of rutin conversion products after incubation with rutin. Consistent with this, there was substantial variability in the abundance of rutin-responsive microbiota among different individuals. Specifically, we observed that Enterobacteriaceae were associated with conversion of rutin into quercetin-3-glucoside and Lachnospiraceae were associated with quercetin production. This suggests that individual microbiotas differ in their ability to metabolize rutin and utilize different conversion pathways.

Contribution to the field

Rutin is a flavonol present in many fruits and vegetables. Rutin and its conversion products exert a wide range of benefits to human health such as anti-antioxidant, anti-cancer, anti-hypercholesterolemia, anti-diabetic, anti-aging, anti-hypertensive activities. The human intestine is inhabited with trillions of microorganisms that play crucial roles in many physiological functions such as protection against pathogenic bacteria, modulation of the immune system, production of vitamins, and fermentation of indigestible plant polysaccharides. Human gut bacteria are able to metabolize many compounds, including flavonols. As the role of the gut microbiota in rutin metabolism has not been systematically evaluated, we investigated rutin conversion by gut bacteria in healthy participants. We find marked inter-individual variability in rutin transformation, and propose a core rutin-stimulated microbiota implicated in rutin transformation. Our findings present new insights into rutin metabolism in healthy humans, which will be helpful for future studies on flavonol metabolism in health as well as in disease conditions.

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Studies involving animal subjects
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Studies involving human subjects
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Inclusion of identifiable human data
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Data availability statement

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Alessandra Riva¹, Ditta Kolimár², Andreas Spittler³, Lukas Wisgrill⁴, Craig W. Herbold¹, László Abrankó² and David Berry¹,⁵*

¹Centre for Microbiology and Environmental Systems Science, Department of Microbiology and Ecosystem Science, Division of Microbial Ecology, University of Vienna, Vienna, Austria.
²Faculty of Food Science, Department of Applied Chemistry, Szent István University, Budapest
³Core Facility Flow Cytometry & Department of Surgery, Research Lab, Medical University of Vienna, Vienna, Austria
⁴Division of Neonatology, Pediatric Intensive Care and Neuropediatrics, Department of Pediatrics and Adolescent Medicine, Medical University of Vienna, Vienna, Austria.
⁵Joint Microbiome Facility of the Medical University of Vienna and the University of Vienna, Vienna, Austria.

*Corresponding author:
David Berry
Althanstrasse 14, 1090 University of Vienna, Austria
Phone: +43 1 4277 76612
david.berry@univie.ac.at

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In review
Abstract

The gut microbiota plays a pivotal role in the conversion of dietary flavonoids, which can affect their bioavailability and bioactivity and thereby their health-promoting properties. The ability of flavonoids to stimulate the activity of the microbiota has, however, not been systematically evaluated. In the present study, we used a fluorescence-based single-cell activity measure [biorthogonal non-canonical amino acid-tagging (BONCAT)] combined with fluorescence-activated cell-sorting (FACS) to determine which microorganisms are stimulated by the flavonoid rutin. We performed anaerobic incubations of human fecal microbiota amended with rutin and in the presence of the cellular activity marker L-Azidohomoalanine (AHA) to detect rutin-stimulated cells. We found that 7% of cells in the gut microbiota were active after a 6 h incubation and 23% after 24 h. We then sorted BONCAT-positive cells and observed an enrichment of Lachnospiraceae (Lachnoclostridium, and Eisenbergiella), Enterobacteriaceae, Tannerellaceae and Erysipelotrichaceae species in the rutin-responsive fraction of the microbiota. There was marked inter-individual variability in the appearance of rutin conversion products after incubation with rutin. Consistent with this, there was substantial variability in the abundance of rutin-responsive microbiota among different individuals. Specifically, we observed that Enterobacteriaceae were associated with conversion of rutin into quercetin-3-glucoside and Lachnospiraceae were associated with quercetin production. This suggests that individual microbiotas differ in their ability to metabolize rutin and utilize different conversion pathways.

Introduction

Flavonoids are a group of bioactive polyphenolic compounds present in a wide variety of plant-based foodstuffs. Rutin (quercetin-3-O-rutinoside) is a flavonol glycoside composed of quercetin and rutinose, a disaccharide of rhamnose and glucose. Dietary sources of rutin include tea, green asparagus, onions, buckwheat, wine, eucalyptus, apples, as well as berries (de Araujo et al., 2013; Kumar and Pandey, 2013; Amaretti et al., 2015). Rutin has been shown to have anti-oxidant properties (Ghorbani, 2017) and to exert anti-aging effects on human dermal fibroblasts and human skin (Choi et al., 2016). It also has anti-neurodegenerative properties (Enogieru et al., 2018) and exhibits protective effects against hyperglycemia, dyslipidemia, liver damage, and cardiovascular disorders (Ghorbani, 2017). Additionally, the rutin degradation products quercetin-3-glucoside and quercetin have been found to have anti-inflammatory, anti-oxidant and anti-mutagenic properties (Gibellini et al., 2011; Kumar and Pandey, 2013; Hobbs et al., 2018). Quercetin-3-glucoside also possesses anti-hypotensive, hypolipidemic effects (Gibellini et al., 2011; Kumar and Pandey, 2013; Hobbs et al., 2018) and quercetin has been reported to ameliorate atherosclerosis and dyslipidemia (Salvamani et al., 2014).

The bioavailability of these dietary flavonoids depends on intestinal absorption, which is determined by their chemical composition and, in particular, by the nature of glycosylation (Matsumoto et al., 2004). The glyco-conjugates of quercetin are poorly absorbed in the upper intestinal tract and accumulate in the large intestine. In the colon, members of the gut microbiota can hydrolyze rutin or other glyco-conjugates, removing the sugar moiety and permitting the absorption of the aglycone (Cardona et al., 2013; Amaretti et al., 2015). Therefore, the colonic microbiota is responsible for the extensive breakdown of the original flavonoid structures into low-molecular-weight phenolic metabolites (Cardona et al., 2013). Currently, it is estimated that 500–1000 different microbial species inhabit the gastrointestinal tract, reaching the highest concentration in the colon (up to 10^12 cells per gram of faeces) (Thursby and Juge, 2017). Bacteria that metabolize rutin possess α-rhamnosidases that transform rutin into quercetin-3-glucoside and/or β-glucosidases that either convert quercetin-3-glucoside into quercetin (Braune and Blaut, 2016) or convert rutin directly into quercetin (Olthof et al., 2003). A limited number of bacteria have so-far been shown to have rutin-metabolizing capabilities in pure culture. α-rhamnosidases involved in deglycosylation of flavonoids have been
characterized in Lactobacillus acidophilus, Lactobacillus plantarum (Beekwilder et al., 2009) and Bifidobacterium dentium (Bang et al., 2015). The capability to degrade rutin into quercetin was reported for Bacteroides uniformis, Bacteroides ovatus (Bokkenheuser et al., 1987), and Enterobacterium avium (Shin et al., 2016). Parabacteroides distasonis was shown produce both quercetin-3-glucoside and quercetin via α-rhamnosidase and β-glucosidase activity (Bokkenheuser et al., 1987), and Eubacterium ramulus and Enterococcus casselilavus are able to convert quercetin-3-glucoside in quercetin (Schneider et al., 1999).

Previous studies of rutin conversion by gut bacteria have involved screening strain collections, which gives limited insight into identifying which bacteria are actually involved in metabolizing rutin in the complex gut microbial community. In the present study, we identified rutin-stimulated cells in the gut microbiota by performing anaerobic incubations of human fecal microbiota amended with rutin in the presence of the cellular activity marker L-Azidohomoalanine (AHA). By sorting active cells and profiling active and total communities using 16S rRNA gene amplicon sequencing we were able to identify specific taxa enriched in rutin-treated samples. We observed marked inter-individual variability in both the extent of rutin degradation product formation as well as the abundance of the rutin-responsive microbial community. Our findings present new insights into rutin metabolism by different microorganisms in healthy individuals, which will be useful for future studies on flavonol metabolism in health as well as in disease conditions.

Material and methods

Sample collection

Fresh faecal samples were collected from 10 healthy subjects (7 females and 3 males, age mean±SD: 30.5±5.8; BMI:mean±SD: 22.19±2.9). All participant followed an omnivore diet. Participants with antibiotic, probiotic, or prebiotic usage in the previous six months were excluded. The study was approved by, and conducted in accordance, with the University of Vienna ethics committee (Reference number 00161) and written informed consent was signed by all enrolled participants.

Anaerobic incubations

Fresh stool samples were immediately introduced into an anaerobic tent. Phosphate-buffered saline (PBS) was added to the sample to arrive to a concentration of 1 g/10 ml. The suspension was homogenized by vigorous shaking and vortexing. Samples were left for 15 minutes to allow large particles to settle and subsequently serially-diluted 1:10 twice. Samples were incubated in autoclaved Hungate tubes in the presence of 1 mM of the non-canonical amino acid L-azidohomoalanine (AHA) (baseclic GmbH, Germany) and 500µM rutin dissolved in dimethyl sulfoxide (DMSO; Sigma-Aldrich). A negative control containing DMSO and a positive control with 2 mg/ml of glucose were used for each experiment. An abiotic control for each time point was included to assess the chemical stability of rutin under the incubation conditions. Samples were incubated under anaerobic condition with a final volume of 5 ml for 0, 6 or 24 hours. Subsequently, samples were centrifuged at 14,000 rpm for 10 min and the supernatant was collected and diluted with equal volume of pure acetonitrile (ACN) in order to stabilize the supernatant samples and then stored at -20°C for LC-HRMS analysis. Part of the sample was frozen for nucleic acid extraction and part was washed twice in PBS and then fixed in 1:1 ethanol:PBS for FACS sorting.

Liquid chromatography-high resolution mass spectrometry (LC-HRMS)

Acetonitrile (LC-MS grade) and formic acid were purchased from VWR. High purity water (18.2 MΩ cm⁻¹) was used for dilution of samples and the preparation of mobile phases (Milli-Q Synergy/Elix water purification system, Merck). Authentic reference standards of rutin and quercetin were purchased from Sigma-Aldrich (> 94%, HPLC) and Extrasynthese (> 99%, HPLC) respectively. Supernatants from the incubations were immediately centrifuged and aliquots for HPLC analysis was
removed and equal volume of CAN was added. Samples were kept frozen until analysis. The ACN-stabilized samples were thawed and homogenized by vortexing. 100 µl was diluted 1:4 in water to decrease acetonitrile content to 10%. The diluted sample was filtered through a 0.22 µm pore nylon mesh syringe filter (Cronus, LabHut Ltd.), and 5 µL were injected into the LC system. Chromatographic separation was achieved on a Phenomenex Kinetex EVO C18 100 x 2.1 mm, 2.6 µm column utilizing an Agilent 1200 HPLC system. The column was operated at 30°C. The binary mobile phase consisted of H2O with 0.1% formic acid (eluent A) and acetonitrile (eluent B). The flow rate was set to 0.4 mL min⁻¹. Gradient separation was started at 5% B and linearly increased to reach 90% in 9 min. The eluent was kept constant at 90% B until 11.5 min and then the column was re-equilibrated at the initial conditions for 11.5 min. The effluent of the LC system was connected to an Agilent 6530 high-resolution, accurate mass quadrupole/time-of-flight mass spectrometer equipped with a dual sprayer electrospray ion source (ESI-Q/TOFMS). The mass spectrometry was run in full scan (MS-only) mode scanning from m/z 50-1700 in negative ionization mode. A continuous reference mass correction was applied using purine and HP-921 (Hexakis(1H,1H,3H-perfluoropropoxy)phosphazene) as reference substances. The ion source temperature was maintained at 325 °C and capillary and fragmentor voltages were set to -4000 V and 140 V, respectively. The Mass Hunter Workstation software package (B02.01) was used for data acquisition and data evaluation.

**BONCAT labelling of microbial cells**

Cu(I)-catalyzed click labelling of chemically-fixed microbial cells was performed on slides as described previously (Hatzenpichler et al., 2014). Briefly, fixed samples were immobilized on glass slides, dried in a 46°C hybridization oven, and dehydrated and permeabilized by placing slides for 3 min sequentially in 50, 80 and 96% ethanol. Then, 1.25 µl of 20 mM CuSO4, 2.50 µl of 50 mM tris[(1-hydroxypropyl-1H-1,2,3-triazol-4-yl)methyl]amine (THPTA) (baseclick GmbH, Germany), and 0.30 µl of alkyn dye (in DMSO) (Jena Bioscience, Germany) were mixed and allowed to react for 3 min at room temperature (RT) in the dark. In the meantime, 12.5 µl of freshly-prepared 100 mM sodium ascorbate (Sigma-Aldrich) and 12.5 µl of 100 mM aminoguanidine hydrochloride (Sigma-Aldrich) were added to 221 µl 1× PBS (pH 7.4). Then, the dye premix was added to this solution, the tube inverted once and samples were covered by 30 µl of solution. Slides were transferred into a humid chamber and incubated in the dark at RT for 30 min. Afterwards, slides were washed three times for 3 min each in 1× PBS and then treated with an increasing ethanol series (3 min each in 50, 80 and 96% ethanol) and air-dried (Hatzenpichler et al., 2014). 1:1000 DNA stain, 4′, 6 diamidino-2-phenylindole (DAPI) solution, (in PBS) was applied for 5 min and then slides were washed in cold MILLI-Q water (Millipore GmbH, Vienna, Austria). Samples were embedded with CitiFluor (Agar Scientific Ltd., Stansted, UK) if used immediately or stored at -20°C. Representative BONCAT pictures of a faecal sample incubated with rutin at 6 and 24h and a negative control containing dimethyl sulfoxide (DMSO) are shown in **Supplementary Figure 1**. For FACS sorting, in-solution click labelling was performed immediately before FACS sorting. For click labelling, 300-500 µl fixed samples were centrifuged at 10,000 rpm for 10 min and re-suspended in 96% ethanol. The pellet was left for 3 min at RT, then centrifuged 10,000 rpm for 5 min. A master mix containing the dye solution was prepared as described above. Samples were suspended in 60-100 µl of solution and incubated in the dark at RT for 30 min. Afterwards, samples were washed three times by centrifugation with 1x PBS (Hatzenpichler et al., 2014). Immediately before sorting, samples were filtered with a 35µm nylon mesh using BD tubes 12x75mm (BD, Germany).

**Image acquisition and analysis**

20-30 images were collected for each sample with an epifluoresce microscope (Zeiss-Axio-imager, Germany). Image analysis was performed using the software *digital image analysis in microbial ecology* (Daim) and the biovolume fraction, the fraction of BONCAT-labelled biomass (Cy5-labelled) relative to the total biomass (DAPI-labelled), was calculated (Daims et al., 2006).
Fluorescence activated cell sorting (FACS)
For flow cytometry sorting, bacteria were labeled in Cy5 dye as previously described, and analyzed on an ultra high-speed cell sorter MoFlo Astrios EQ (Beckman Coulter, Brea, CA, USA) using the Summit v6.2 software (Beckman Coulter). To standardize the daily measurement and to assess the size of the bacteria, calibration beads (silica beads 100, 500 and 1000 nm, Kisker Biotech, Steinfurt, Germany) having a refractive index close to biological material were recorded. The sorting of Cy5-labeled bacteria was performed as follows: In a first scatter plot, the 561nm SSC Height-Log parameter was set vs. the 488nm FSC1 Height-Log parameter. To reduce electronic noise the triggering signal was set on the 561nm SSC parameter. A second dot plot 488nm FSC1-Height-Log vs. 488nm SSC-Height-Log showed in a first measurement the different sizes of the silica beads and in the following measurements the scattering of the bacteria. Bacteria were then pre-gated and displayed on a third scatter plot with 488nm SSC area log axes vs. 640nm 671/30-Area-Log axes. Cy5-positive bacteria were then sorted out into tubes with a maximum event rate of 50,000 events per second. Reanalysis of the samples showed a purity of >99%. A overview of the gating strategy and FACS selectivity analysis are shown in Supplementary Figure 2 and 3.

DNA extraction and 16S rRNA gene amplicon sequencing
DNA extraction was performed for both total microbial community and the FACS-sorted fraction using the QiAmp mini DNA extraction kit (Qiagen) according to the manufacturer’s instructions. PCR amplification was performed with a two-step barcoding approach according to Herbold et al., 2015 using 16S rRNA gene primers targeting most bacteria (S-D-Bact-0341-b-S-17 [5’-CCTACGGGNGGCWGCAG-3’]  and S-D-Bact-0785-a-A-21 [5’-GACTACHVGGGTATCTAATCC-3’]). The barcoded amplicons were purified with ZR-96 DNA Clean-up Kit (Zymo Research, USA) and quantified using the Quant-iT™PicoGreen® dsDNA Assay (Invitrogen, USA) (Herbold et al., 2015). An equimolar library was constructed by pooling samples, and the resulting library was sent for sequencing on the Illumina MiSeq platform at Microsynth AG (Baligach, Switzerland).

Sequence processing and data analysis
16S rRNA gene sequence data were sorted into libraries according to Herbold et al. (2015) and processed into amplicon sequence variants (ASVs) using the Divisive Amplicon Denoising Algorithm (DADA2) (Callahan et al., 2016) and classified using the RDP classifier (Wang et al., 2007) as implemented in Mothur (Schloss et al., 2009). Sequencing libraries were subsampled to a smaller number of reads than the smallest library (subsampling to 1000 reads) to avoid biases related to uneven library depth. 16S rRNA gene sequence data has been deposited in the NCBI Short Read Archive under PRJNA622517. Statistical analysis was performed using R statistical software (https://www.r-project.org/). Statistical analysis to compare samples groups was performed using ANOVA, and with the R package DEseq2 (Love et al., 2014). The statistical significance of factors affecting microbiota composition was evaluated using non-parametric permutational multivariate analysis of variance (perMANOVA), significant clustering of groups was evaluated with analysis of similarities (ANOSIM), ordination was performed using redundancy analysis (RDA) and non-metric multidimensional scaling (NMDS) in the vegan package in R (Oksanen et al., 2010). Alpha and beta diversity metrics were also calculated with the vegan package. Variables are expressed as mean ± SD (standard deviation). A probability value (p-value) less than 0.05 was considered statistically significant and p-values were adjusted with the False Discovery Rate method (FDR) in the case of multiple comparison. Statistical analysis to compare producer groups was performed using ANOVA and Tukey test for multiple comparisons.
Results

Biotransformation of rutin by the gut microbiota

In order to characterize the biotransformation of rutin by the gut microbiota and to identify bacteria stimulated by rutin, we performed anaerobic incubations of freshly-collected stool contents amended with 500µM rutin (Figure 1). This concentration is in line with previous studies and is consistent with a reasonable dietary intake (Amaretti et al., 2015; Zamora-Ros et al., 2016). Over the course of the incubations there was a slight but not statistically-significant reduction in rutin (one-way ANOVA, p = 0.41, n = 10 Figure 2A). However, Q-gluc (quercetin-3-glucoside) and Q (quercetin) appeared after incubation with biomass (p = 0.019 and 0.036 for Q-gluc and Q, respectively, n = 10; Figures 2B and C), indicating that rutin was actively transformed by the gut microbiota. Low levels of Q-gluc were present in all time zero samples (Figure 2B), which may be because rutin preparations were either not completely purified or chemically degraded to Q-gluc during storage. Interestingly, there was substantial variation in the amount of Q-gluc and Q formed during incubation with the gut microbiota, suggesting considerable inter-individual variability in the capacity to metabolize rutin by different gut microbial communities (Figure 2D and E).

The core rutin-stimulated microbiota

As the capacity of dietary polyphenols to modulate the activity and/or composition of the gut microbiota is still poorly understood, we aimed to identify the microbial taxa stimulated by rutin. Rutin amendment stimulated a subset of the microbiota, and active cells were detected in almost all samples after 6 h of incubation with rutin, with an increasing number after 24 h (6h: 7.3 ± 7.0%, 24h: 29 ± 11.7% [mean ±SD]; t-test: p = 0.0003, n = 20) (Figure 3A,B). The diversity and the composition of the total microbial community did not change significantly during the short-term incubations (Figures 4A,B; perMANOVA, p = 0.99), indicating that the applied incubation conditions did not appreciably modify the composition of the microbiota (i.e. there was no strong “bottle effect”). However, the rutin-stimulated fraction of the community, as determined by BONCAT activity labeling and FACS sorting, was significantly different from the total community. The diversity of the active fraction was lower than the total community (ANOVA, p<0.0001 for all tested alpha diversity metrics; Figure 4A), and the NDMS ordination showed a clear separation of samples between the total community and the active fraction (perMANOVA, p < 0.0001; ANOSIM, p = 0.001) (Figure 4B). The most abundant taxa detected in the active fraction were Bacteroidaceae, Enterobacteriaceae, Lachnospiraceae, Tannerellaceae, and Ruminococcaceae (Figure 5). We found that members of the Enterobacteriaceae (Escherichia/Shigella), Tannerellaceae (Parabacteroides), Erysipelotrichaceae (Erysipelotrichales), and Lachnospiraceae (Lachnolocustrium and Eisenbergiella) were significantly increased in the active fraction compared to their respective total community samples (negative binomial distribution, Wald test, p<0.05, n=20).

Microbial community composition is associated with rutin transformation patterns

We observed that there was a large variability in the ASVs that were enriched in the active fraction of the community across the incubations from different donors. Only 13/97 and 23/91 ASVs were shared among all incubations at 6 h and 24 h, respectively (Supplementary Figure 4A,B). Consistent with this, we found that the microbial community varied significantly within participant in both the total community and active fraction (perMANOVA: p <0.0001, Supplementary Figure 4C,D), which was in line with the observation that different cell morphologies were observed in the active fraction from different donors (Figure 3B). Though stool incubations from all participants led to the production of rutin degradation products, the pattern of rutin product formation was markedly different among different participants. We therefore divided the samples into the following groups: "High Q-gluc producers", "High Q producers", and "Low producers". Based on these categories, we observed a significant clustering of both total community and active fraction samples in the redundancy analysis ordination (perMANOVA, total community: p = 0.001, active fraction: p =
In the active fraction, we observed an enrichment of Enterobacteriaceae (Escherichia-Shigella) in High Q-glc producers, and this taxon represented the majority of sequences recovered in the active fraction by 24 h (ANOVA: 0.0002, n=50). In contrast, High Q producers had an enrichment of Lachnospiraceae (Lachnoclostridium and Eisenbergiella), which was the dominant taxon in the active fraction by 24 h (ANOVA: 0.0257, n=50). Low producers had a trend, though not statistically significant, towards enrichment in Tannerellaceae (Parabacteroides) and Erysipelotrichaceae (Erysipelatoclostridium) in the active fraction (Figure 6C, Supplementary Figure 5).

**Discussion**

The gut microbiota plays a key role in the conversion of dietary flavonoids. Though conversion of flavonoids by the microbiota has gained increasing interest (Braune and Blaut, 2016), key microbial players in flavonoid metabolism remain poorly characterized (Cardona et al., 2013). In this study, we investigated metabolism of the flavonoid rutin by the human gut microbiota. Rutin conversion products were detected in all tested donor stool incubations, but there was a dramatic variation in the amount of Q-glc and Q produced by different microbioso. This suggests inter-individual variability in preference or capability for rutin metabolism. This is in line with a previous report of high person-to-person variation in the concentration of phenolic acids between 2 and 24 h incubation of stool with rutin (Jaganath et al., 2009).

Alpha diversity was lower in the active fraction compared to the total community, implying that a subset of the community is stimulated by rutin. In our study, the significant bacterial taxa selected are: Lachnospiraceae (Lachnoclostridium, Eisenbergiella), Enterobacteriaceae (Escherichia), Tannerellaceae (Parabacteroides), and Erysipelotrichaceae (Erysipelatoclostridium). These bacteria may represent a “core rutin-selected microbiota” in healthy individuals. Members of the Enterobacteriaceae (Escherichia coli, Escherichia fergusonii, and Enterobacter cloacae) have been previously implicated in the O-deglycosylation and dehydroxylation of different flavonoids (Miyake et al., 1997; Hur et al., 2000; Zhao et al., 2014; Braune and Blaut, 2016). In a batch culture fermentation experiment, Tzounis et al. found that catechin, a flavan-3-ol monomers, promoted the growth of E. coli (Tzounis et al., 2008) and Duda-Chodak found that rutin does not inhibit E. coli growth (Duda-Chodak, 2012). In accordance with our findings, Parabacteroides distasonis has been shown to produce both quercetin-3-glucoside and quercetin (Bokkenheuser et al., 1987) and the Lachnospiraceae members Blautia and Eubacterium were able to convert rutin in quercetin (Kim et al., 2014) and quercetin-3-glucoside in quercetin, respectively. Based on rutin degradation capability and the identification of the core-rutin selective microbiota, we divided the donor microbios into high Q-glc, high Q, and low producers. We observed that members of Enterobacteriaceae (Escherichia) were associated Q-glc production, putatively due to expression of alpha-rhamnosidases that can act on rutin. We also found that Lachnospiraceae (Lachnoclostridium and Eisenbergiella) were associated with Q production, which may be due to expression of beta-rutosidase enzymes or a combination of alpha-rhamnosidase and beta-glucosidase enzymes.

According to our findings, we hypothesize that inter-individual variability in rutin metabolism is driven by differences in the composition of the gut microbiota. Variation in rutin and other flavonoid metabolisms in humans may also be caused by host and environmental factors as diet and genetic polymorphism and differences in enzymatic activity (Almeida et al., 2018). In conclusion, individual microbioss exert distinct capability in rutin utilization, showing a higher response in certain individuals, whereas others seem less capable in rutin utilization. Future research that takes into account functional gene analysis, diet, and host physiology will advance our understanding of the role of the gut microbiota in rutin degradation and provide opportunities to improve human health.
Abbreviations: Q-glc, Quercetin-3-glucoside; Q, Quercetin; FACS, fluorescence activated cell sorting.

Ethics statement
The study was approved by, and conducted in accordance, with the University of Vienna ethics committee (Reference number 00161) and written informed consent was signed by all enrolled participants.

Data availability statement
16S rRNA gene sequence data has been deposited in the NCBI Short Read Archive under accession number PRJNA622517.

Author contributions
DB, LA and AR conceived and designed the experiments. AR performed the experiments and data analyses. AR and DK performed anaerobic incubation experiments. AS, LW and AR performed FACS sorting. CH performed bioinformatics analyses. AR and DB wrote the paper. All authors have given approval to the final version of the manuscript.

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Conflict of interest
The authors declare no conflict of interest.

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**Figure legends:**

**Figure 1.** Schematic representation of rutin degradation. Rutin is present in a wide variety of foods. Rutin is not well-absorbed in the small intestine of humans, and thus is transported into the colon and metabolized by the gut microbiota into quercetin-3-glucoside and then quercetin or directly into quercetin. Quercetin may be subsequently degraded mainly into different phenolic acids.

**Figure 2.** Levels of rutin, quercetin-3-glucoside (Q-glc) and quercetin (Q) after 0, 6 and 24 h incubation. (A-C) Box and whisker plots of the combined results of incubation of rutin with the gut microbiota of 10 participants, shown as peak areas obtained from LC-HRMS analysis. Open circles indicate results of abiotic controls. (D) Q-glc and (E) Q levels in each incubation after 6h and 24h.

**Figure 3.** The rutin-stimulated microbiota. (A) Relative abundance of rutin-stimulated cells in incubations. (B) Representative microscopic images of samples from three participants showing variability in cell morphology between donors. Active cells are represented in red (BONCAT-Cy5) and all cells are stained in blue (DAPI).
**Figure 4.** Microbiota richness and diversity in rutin stimulated samples. (A) Observed ASVs, Chao1 estimated richness, Shannon diversity, and inverse Simpson diversity estimators show significant difference between time points, active fraction and total community [ASVs richness, Chao1 and Shannon (ANOVA: p<0.0001, n=50), Inv. Simpson (ANOVA: p=0.020, n=50)]. Multiple comparisons are represented in the figure as asterisks. (B) NMDS ordination shows samples separation between the total community and the active fraction.

**Figure 5.** Relative abundance of bacterial taxa based on 16S rRNA gene amplicon sequencing at family and genus level for each participant. (A, B) Relative abundance of the total community and the active fraction at time 6 and 24 h. (C, D) Relative abundance of the active fraction at time 6 or 24 h. Family and genera with relative abundance >0.5% and >1%, respectively, is shown.

**Figure 6.** Samples clusters based on rutin degradation pattern. Redundancy analysis shows sample clustering by degradation pattern in both the (A) total community (constrained variance explained:18.6%, RDA1 (14%), RDA2 (4.8%)) and (B) active fraction (constrained variance explained: 40.5%, RDA1 (36.3%), RDA2 (4.1%)) (C) Family-level heatmap showing the square root transformed relative abundance of the active fraction divided by producer groups at 24h. Significant changes in relative abundance (Low producers vs. High producers) are indicated with asterisks. *Enterobacteriaceae* increased in relative abundance in the High Q-glc producers. (High Q-glc producers vs. Low producers, p= 0.0055, High Q-glc producers vs. High-Q producers, p= 0.0061). *Lachnospiraceae* increased in relative abundance in the high-Q producers (High-Q producers vs. Low producers, p=0.0043, High-Q producers vs. High Q-glc producers, p<0.00001).
Dietary rutin: asparagus, onions, apples, berries, wine and tea, buckwheat, eucalyptus

Large intestine

Rutin (quercetin-3-rutinose)

α-hamnosidase
β-rutinosidase

Quercetin-3-glucoside

β-glucosidase

Quercetin

Phenolic acids
/review
In review.
Figure 4: TIFF

A

| ASVs richness | Chao1 | Shannon | Inverse Simpson |
|---------------|-------|---------|----------------|
| 150           | 350   | 4       | 3              |
| 120           | 300   | 3.5     | 2.5            |
| 90            | 250   | 3.2     | 2              |
| 60            | 200   | 2.8     | 1.5            |
| 30            | 150   | 2.5     | 1              |
| 1              | 100   | 2       | 0.5            |

B

Active fraction

Total community

Active fraction 0h

Total community 0h

Total community 6h

Total community 24h

Active fraction 6h

Active fraction 24h

NMDS1

NMDS2
Figure 5.TIFF

A

B

C

D

Relative abundance (%)

0 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15

Active fraction

Total community

Active fraction

Relative abundance (%)

0 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15

Active fraction

Total community

Active fraction
