Substrate-Dependent Fermentation of Bamboo in Giant Panda Gut Microbiomes: Leaf Primarily to Ethanol and Pith to Lactate

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The giant panda is known worldwide for having successfully moved to a diet almost exclusively based on bamboo. Provided that no lignocellulose-degrading enzyme was detected in panda’s genome, bamboo digestion is believed to depend on its gut microbiome. However, pandas retain the digestive system of a carnivore, with retention times of maximum 12 h. Cultivation of their unique gut microbiome under controlled laboratory conditions may be a valid tool to understand giant pandas’ dietary habits, and provide valuable insights about what component of lignocellulose may be metabolized. Here, we collected gut microbiomes from fresh fecal samples of a giant panda (either entirely green or yellow stools) and supplied them with green leaves or yellow pith (i.e., the peeled stem). Microbial community composition was substrate dependent, and resulted in markedly different fermentation profiles, with yellow pith fermented to lactate and green leaves to lactate, acetate and ethanol, the latter to strikingly high concentrations (~3%, v:v, within 3.5 h). Microbial metaproteins pointed to hemicellulose rather than cellulose degradation. The alpha-amylase from the giant panda (EC 3.2.1.1) was the predominant identified metaprotein, particularly in reactors inoculated with pellets derived from fecal samples (up to 60%). Gut microbiomes assemblage was most prominently impacted by the change in substrate (either leaf or pith). Removal of soluble organics from inocula to force lignocellulose degradation significantly enriched Bacteroides (in green leaf) and Escherichia/Shigella (in yellow pith). Overall, different substrates (either leaf or pith) markedly shaped gut microbiome assembles and fermentation profiles. The biochemical profile of fermentation products may be an underestimated factor contributing to explain the peculiar dietary behavior of giant pandas, and should be implemented in large scale studies together with short-term lab-scale cultivation of gut microbiomes.

Keywords: giant panda, cellulose, hemicellulose, alpha amylase, lignocellulose, fermentation, gut microbiome, ethanol
INTRODUCTION

The iconic giant panda (*Ailuropoda melanoleuca*) is known worldwide for being a carnivore (belonging to the family of the *Ursidae*, thus a bear) that moved to a vegetarian diet based almost exclusively on bamboo, a member of the grass family *Poaceae* [99% in the wild (Schaller et al., 1985)]. Paleontological and molecular evidence suggest this dietary shift initiated about 2 million years ago (Jin et al., 2007; Zhao et al., 2010). While dentition, skull and jaw musculature adapted to crush and grind the hard bamboo culms, surprisingly the soft tissues of the gut remained those typical of its carnivorous ancestors (Wildt et al., 2006). The latter is considered the main reason for the short retention time of ingested bamboo in the gastrointestinal tract (Dierenfeld et al., 1982), which has no apparent special compartment to retain food contrary to monogastric herbivores and ruminants. Clearance of the digesta is generally observed between 5 and 12 h (Dierenfeld et al., 1982; Mainka et al., 1989; Edwards et al., 2006; Senshu et al., 2007), although longer times have been reported in pandas with intestinal disorders (Mainka et al., 1989; Senshu et al., 2007) or depending on the portion of bamboo ingested [about 8 h for shoots, 10 h for stems and 14 h for leaves (Schaller et al., 1985)]. The low digestibility of bamboo by pandas [<40% on either a dry matter or apparent energy conversion basis (Dierenfeld et al., 1982; Senshu et al., 2007; Sims et al., 2007; Finley et al., 2011)], raised some questions over the evolutionary advantage of moving to such a diet (Gittleman, 1994). How giant pandas meet their nutritional and energetic requirements has not been fully resolved, particularly when compared to other large herbivores. Recent reports suggest that giant pandas are macronutritional carnivores (Nie et al., 2019), with the low efficiency of bamboo degradation compensated by maintaining a high daily feed intake [6 to 15% of body weight dry matter (Mainka et al., 1989; Dierenfeld, 1997)], constraining pandas to spend up to one third of their day in feeding (Edwards et al., 2006). In this context, short gut retention times and low energy expenditure of daily giant panda’s life are considered an assumption that the panda’s gut microbiome may fulfill this function. The essential contribution of gut microbiomes in lignocellulose degradation has been extensively reported in nature, in mammals [e.g., historically in ruminants (Gordon and Phillips, 1998); and recently more in particular in monkeys (Xu et al., 2015); beavers and mooses (Wong et al., 2016); wallabies (Pope et al., 2010); reindeers (Pope et al., 2012), annelids [e.g., earthworms (Fujii et al., 2012)], and insects [e.g., beetles (Geib et al., 2010; Huang et al., 2012; Sari, 2016)]; cockroaches (Bertino-Grimaldi et al., 2013); and most extensively in termites (Warnecke et al., 2007; Mathew et al., 2013; Brune, 2014; Do et al., 2014)]. Although not representative of the entire gastrointestinal tract, gut microbiomes in giant pandas are generally inferred using microbial communities from stool samples. Such microbial communities have been investigated for their assemblage (Zhu et al., 2011), how they related to dietary shifts (Sims et al., 2007; Williams et al., 2013; Knott et al., 2017) and for their potential cellulose (Zhu et al., 2011), hemicellulose (Zhang et al., 2018) or lignin biodegradation capacity (Fang et al., 2012), as assessed by metagenome analysis. However, no attempt was made so far to cultivate gut microbiomes to mimic giant panda’s feeding. In the present study, we collected the fresh fecal material generated by a young, male panda and used it as microbial inoculum in lab-scale tests. The objective was to put the emphasis on the products of fermentation rather than on its substrates (i.e., the different portions of bamboo). While a relation to the chemical composition of the ‘substrates’ has been made (i.e., the different portions of ingested bamboo), whether panda’s dietary choices also relate to the ‘product’ of bamboo fermentation has been neglected thus far. Here, we focused on the interplay between fermentation of different portions of bamboo of the species *Phyllostachys bissetii* (either the green leaf or the yellow pith), the profile of short-chain organics and biogas produced, the gut microbiome community assemblage (originated from stools either entirely green or yellow) and the microbial metabolism as assessed by metaproteomics.

RESULTS

Characterization of Feces and Bamboo

Entirely green or yellow stools from a young, male panda were characterized (Figure 1). Green and yellow stools were typically populated by about 1.0–1.5 × 10^9 microbial cells g^-1 dry weight (DW) matter. In green stools, the profile of short-chain organics was a heterogeneous mix of lactate, acetate and...
Ethanol (900 to 1400 mg g\(^{-1}\) DW), with minor concentration of formate, contrary to yellow stools where lactate was the only abundant compound (4000 mg g\(^{-1}\) DW). *P. bisetti* is a bamboo species commonly offered to pandas in captivity (Edwards et al., 2006). In this study, *P. bisetti* had the typical biochemical composition for the autumn-winter period, i.e., when it was collected (Supplementary Table S1): both leaves and piths have rather high levels of cellulose, hemicellulose and lignin, however, hemicellulose is typically higher in leaves (about 30%) and lignin in piths (about 15%) (Tabet et al., 2004). Predominant genera (Figure 1B) and microbial community assemblage (as assessed by PCoA analysis, Figure 1D) were highly similar in either green leaf or yellow pith of the bamboo species *P. bisetti*, however, they differed remarkably in the green and yellow stools (Supplementary Table S2). This indicates that bamboo passage through the panda’s gut results in enrichment of specific microbial members, organized in a markedly distinct manner as compared to those natively present on the ingested bamboo.

**Effect of Selected Portions of Bamboo and Fecal Microbial Communities on Fermentation**

The fermentation capacity of different portions of *P. bisetti* by giant panda fecal microbial communities was investigated. A ‘green fermentation line’ was set up using bamboo leaf as substrate, and entirely green fecal samples as starting material to mimic giant panda’s gut microbiomes. Similarly, a ‘yellow fermentation line’ was prepared using the yellow pith, and entirely yellow fecal samples (preparation protocol summarized in Figure 2). The green and yellow bacterial inocula thus prepared carried some soluble organics derived from...
the actual giant panda’s gut digestion system. Therefore, two other conditions were tested in parallel (one per fermentation line), where the freshly prepared inocula were subjected to centrifugation to remove the soluble organics (Figure 2). The pellet generated by centrifugation contained bacterial biomass along with small fragments of non-digested bamboo; it was re-suspended in a phosphate buffer saline (PBS) solution and provided with fresh bamboo (either leaf or pith). Therefore, accumulation of fermentative products in such de-watered (de-H$_2$O) inocula did not depend on soluble organics previously generated in the panda’s gut rather on the fresh, solid bamboo supplied at the beginning of the experiment.

Gas production (either H$_2$ or CO$_2$) was significantly higher in the green fermentation line as compared to the yellow (log$_2$ fold change $\log_2^{fc} + 4.61, p = 0.007$; Figures 3A,B). In particular, H$_2$ was produced in very low titers when supplying the yellow
pith. In all conditions, gas production rates peaked at 3.5 h of experiment and declined thereafter (Figures 3C,D). Only in the green fermentation line gas production was still substantial after 14 h, which is the longest observed retention time in the giant panda’s gut under normal physiological conditions. The removal of soluble organics tested with de-H2O inocula reduced H2 and CO2 production, with the highly producing green fermentation line the most impacted. H2 consumption initiated after 27 h in all cultures that had previously accumulated it (Figure 3C). CH4 was never detected in any condition (detection limit 0.01%) demonstrating a purely fermentative metabolism by panda’s fecal microbial communities.

The pH of the two fermentation lines differed markedly (p < 0.00001), with the green line around 6.0 and the yellow around 4.5 (Figure 3E). Resuspension in PBS after the dewatering process had little effect on pH in either line despite the initial pH of PBS (which was 7.35).

The green fermentation setup resulted in the accumulation of a variety of organics contrary to the yellow, which was dominated by lactic acid (Figure 4). Such fermentation profiles were highly consistent with what detected in the fecal material they originated from, respectively (Figure 1), indicating that our cultivation system was able to reproducibly mimic the giant panda’s gut conditions. When considering the longest retention time observed in panda’s guts [14 h (Schaller et al., 1985)], the net accumulation of fermentative products was slightly affected in the green line (~30% upon dewatering, compare Figures 4A and C) and not affected at all in the yellow line (compare Figures 4B and D). However, fermentation capacity of de-H2O inocula remained almost unchanged following 14 h of incubation, contrary to full inocula which reached about 2 times higher concentrations over 40 h (Figure 4). Fermentation profiles were not affected by dewatering in either line. Overall, this indicated that gut microbiomes from both green and yellow stools were consistently capable of fermenting solid bamboo, and could generate short-chain organics at almost equivalent rates within panda’s gut retention times.

In the green line, the fermentative products which accumulated most rapidly (within the first 3.5 h, Figure 4) were lactate and ethanol, along with traces of acetate. In
particular, ethanol net production was higher than 1.1 $\text{gCOD L}^{-1}$ (about 250 mg L$^{-1}$ or about 3%, v:v), which accumulated within only 3.5 h, that is, well within the shortest retention time commonly observed in giant pandas [5 h (Dierenfeld et al., 1982)]. This suggests that giant pandas feeding on bamboo leaves may experience high ethanol levels in their gut, in line with ethanol abundance in green steools (Figure 1).

A time-dependent consumption of lactate (starting after 7 h of incubation) was consistent with the production of acetate, propionate and butyrate (Figures 4A,C). Acetate can derive from conversion of acidogenic products (e.g., ethanol oxidation), propionate from lactate reduction, and butyrate from either glucose (or lactate) oxidation, or from acetate reduction. Furthermore, with prolonged retention times the product type shifts from more oxidized compounds, such as acetate or lactate, toward more reduced compounds (Arslan et al., 2016), as in fact observed here with longer fatty acids accumulated more prominently only after 15–20 h. In stark contrast, lactate was steadily produced in the yellow fermentation from 3.5 to 40 h (Figures 4B,D).

In giant panda’s digestibility trials, total solids (TSs) are reduced 7 to 40% within 13 h (Dierenfeld et al., 1982; Mainka et al., 1989; Sims et al., 2007; Finley et al., 2011). The longer incubation time in the present study (40 h) resulted in higher TS reductions, about 50 and 30% in green and yellow full inocula, respectively (Supplementary Figure S1A), with de-$\text{H}_2\text{O}$ inocula showing a TS removal > 60% (Supplementary Figure S1A). The latter confirmed that replacing soluble organics resulting from panda’s digestion with PBS forced gut microbiomes to access solid bamboo. A similar trend was observed with volatile solids (VSs) (Supplementary Figure S1B).

**Microbial Communities**

Green and yellow fermentation lines had a different cell growth during the incubations. The cell number increased significantly in the green line (from 0.67 to $1.95 \times 10^9$ cells mL$^{-1}$ in full inocula $\log_{2}\text{c} + 1.54$, $p = 0.0003$; and from 0.57 to $1.46 \times 10^9$ cells mL$^{-1}$ in full de-$\text{H}_2\text{O}$ inocula, $\log_{2}\text{c} + 1.35$, $p = 0.01$; Figure 5A), with slightly more intact cells at the end of the incubation (about 75%, Figure 5B). On the contrary, in the yellow line no net increase in cell density was observed while intact cells were strongly reduced (to about 25%, Figure 5B).

The diverse spectrum of short-chain organics in the green fermentation line (Figures 4A,C) was mirrored by a highly diverse group of microbial species (Figure 5C), with *Escherichia/Shigella* (OTU00001), *Veillonella* (OTU00002 and OTU00005), *Acidaminococcus* (OTU00012), *Megasphaera* (OTU00023), *Streptococcus* (OTU00003), *Bacteroides* (OTU00008, OTU00010, and OTU00007) and *Clostridium sensu stricto* (OTU00004) above 1% relative abundance (Figure 5E and Supplementary Table S3). Similarly, the accumulation of almost exclusively lactate in yellow inocula (Figures 4B,D) was reflected into communities with low diversity (Figure 5C) where either *Streptococcus* (OTU00003, in Yellow inocula) or *Escherichia/Shigella* (OTU00001, in De-$\text{H}_2\text{O}$ yellow inocula) constituted about 60% of the community (Figure 5E and Supplementary Table S3). A Lactobacillales *Leuconostoc*-related strain (OTU00006) was typically associated with yellow rather than green fermentation lines (Figure 5E) (from 0.2 to 14.7%, respectively, $\log_{2}\text{c} + 6.2$, Supplementary Table S3). Microbial community assembly differed markedly in green and yellow lines, with the removal of soluble organics through dewatering also a factor in shaping communities (Figure 5D). In particular, when forcing communities to feed solely on solid bamboo leaf, the relative abundance of some *Bacteroides* increased (in particular OTU00008 from 0 to 10.4%; and OTU00010 from 0 to 5.3%; Supplementary Table S3) along with *Parabacteroides* (OTU00014, from 0 to 2.0%) and *Morganella* (OTU00019 from 0 to 1.2%, Supplementary Table S3). Concerning the yellow fermentation line, only *Escherichia/Shigella* (OTU00001) increased remarkably when forcing communities to feed solely on solid pith (from 4.5 to 52.8%, $+3.6 \log_{2}\text{c}$ Supplementary Table S3).

**Alpha-Amylase From the Giant Panda Is the Predominant Enzyme in Bamboo Fermentation**

The predominant microbial pathway in any tested condition was glycolysis (between 30 and 50% of all microbial metaproteins; Figure 6 and Supplementary Table S4), with no difference between culture conditions ($p > 0.05$). Glycolysis is the main metabolic pathway processing glucose, a component of the complex heteropolymer hemicellulose (which also contains xylose, arabinose, mannose, and galactose among others) and the only monomer required to build up the linear polymer cellulose (Perez et al., 2002). Metaproteins related to the functions ‘glucose metabolism,’ ‘xylose metabolism,’ and- to a minor extent- ‘arabinose catabolism’ were also detected in all conditions (up to 0.5% of all microbial metaproteins, Figure 6). These metabolic pathways were particularly relevant in the yellow fermentation line when soluble organics were removed, thus forcing gut microbiomes to degrade the yellow pith (i.e., De-$\text{H}_2\text{O}$ yellow inoculum + Pith, Figure 6). Enzymes related to the biological function ‘cellulose degradation’ were < 0.038% of all microbial metaproteins in only two reactors out of 12, and absent in all others. A cellulbiose-specific phosphotransferase enzyme (E.C. 2.7.1.205) suggestive of cellulose degradation was detected, however, it was only found in a few replicates of the yellow fermentation and at very low levels (<0.095% of all microbial metaproteins, Supplementary Table S5). Other highly expressed microbial biological functions (>5% of all microbial metaproteins) were virulence, iron storage, transport and protein biosynthesis (Supplementary Table S4; more results discussed in the Supplemental Information). Metaproteins from yeasts were detected, however, apart from an elongation factor (whose relative abundance was between 1.1 up to 3.1%) the sum of all other metaproteins associated with yeasts was not higher than 0.03% in any of the 12 reactors (Supplementary Table S5).

An analysis of all the metaproteins was conducted including those not taxonomically related to microbial species. Among the carbohydrate active enzymes (CAZy), alpha-amylases were the most represented metaprotein (Figure 7) and almost exclusively assigned to the giant panda (*A. melanoleuca*) (up to 60%
of all identified metaproteins, Supplementary Table S6), with identification of three sequences from Metazoa owing to the high similarity with panda’s alpha-amylases (Supplementary Table S6; all metaproteins in Supplementary Table S7). All identified peptides of alpha-amylase in the giant panda were assigned to the UniProt accession G1L4F3_AILME, although several genes related to alpha-amylase are described in the genome of giant pandas. Loss of material during the dewatering process of the inocula was expected to potentially reduce the amount of water-soluble alpha amylases present in any de-H$_2$O reactor. On the contrary, in either green or yellow fermentation line, alpha-amylases from the panda were far more abundant when soluble organics were removed through dewatering (p < 0.033, log$_2$fc > + 2.68; Figure 7 and Supplementary Table S6), indicating that panda’s alpha-amylases were attached to their solid substrate. Total metaprotein counts were higher in de-H$_2$O reactors at the end of the experiment (+ 0.4 and + 0.7 log$_2$fc in green and yellow line, respectively) excluding a ‘dilution effect’ on relative abundance. The high amount of alpha-amylases in de-H$_2$O reactors suggests that they were equally concentrated in both full and de-H$_2$O inocula at the onset of the incubation, however, their activity may have decreased in full inocula due to the availability of soluble organics (e.g., Figure 1A).

Many other enzymes related to cellulose, hemicellulose, or lignin degradation were detected (Figure 7), however, their concentration was below 0.1% (Supplementary Table S6). Among these, maltase glucoamylases and alpha-mannosidases from the giant panda were significantly more abundant in the green fermentation line when soluble organics were maintained in the inoculum (p < 0.032; Supplementary Table S6). Besides, some uncharacterized proteins of the giant panda were detected. Sequence motifs assigned in UniProt suggest potential protease activity of several hits (Supplementary Table S8). The protein D2HX31_AILME shows a sequence motif related to jacalin-type lectin. Jacalin-related human ZG16p lectin binds oligomeric sugars and is involved in formation of pancreatic zymogen granules as well as in protection against invading pathogen in colon (Kanagawa et al., 2014).

DISCUSSION

Understanding the feeding behavior of the giant panda, the flagship of endangered species with less than 2000 individuals [in the wild and captivity (Huang et al., 2015)] is a critical part of its conservation. Besides, this offers the opportunity to gain knowledge from a unique microbiome, exposed to a high load of lignocellulosic bamboo within a timespan of hours. In the present study, giant panda gut microbiomes were cultivated for the first time under laboratory conditions. Experiments aimed at revealing how microbial community assemblage, fermentation products and metaproteome profiles were influenced after
simulation of giant panda's selectiveness for different bamboo portions. This was attained through either uniquely 'green' or 'yellow' lab-scale fermentations, where both bamboo portions (either leaf or pith, respectively) and their correspondent gut microbiomes were used (Figure 2).

Bamboo digestion followed a distinctive pattern strictly related to the portion of bamboo used as substrate: fermentation of green leaves occurred at a close to neutral pH and generated ethanol, lactate and acetate (Figures 3, 4). The yellow pith occurred at more acidic pH and formed lactate almost exclusively (Figures 3, 4). Such fermentation profiles matched those detected in the stools from which the gut microbiomes originated (Figure 1), indicating that the present setup was representative of the actual fermentative patterns occurring in the giant panda's gut. In particular, in green leaves ethanol rapidly accumulated to strikingly high concentrations (about 3%, v:v). Irrespective of the fermentation type, glycolysis was the predominant carbohydrate-related microbial biological function assessed by metaproteomics, with microbial metaproteins of glucose and xylose metabolism and to a lesser extent- arabinose catabolism detected up to 5% (Figure 6). As glucose, xylose and arabinose are found in hemicellulose, with glucose the only monomer required to build up cellulose (Perez et al., 2002), the present data suggests that the panda's gut microbiomes degraded hemicellulose. Metagenome analysis supports the capacity of gut microbiomes inferred from fecal microbial communities to degrade cellulose (Zhu et al., 2011) and hemicellulose (Zhang et al., 2018), although to a limited degree (Guo et al., 2018). As enzymes related to cellulose digestion were found to very low concentrations or were below detection limits (Supplementary Table S6), our data is in agreement with a limited potential of cellulose degradation in giant panda gut microbiomes. Xylose and arabinose are pentoses, monosaccharides made of five carbon atoms. In heterofermentation, pentoses are used to produce lactate, CO$_2$ and ethanol/acetate mixtures, as opposed to homofermentation where hexoses such as glucose enter glycolysis to generate almost solely lactate (Axelsson and Ahrné, 2000). While showing comparable microbial metaproteomes (Supplementary Table S4), the fermentative product profiles of green leaves typically resembled that of a heterofermentation, as opposed to the yellow pith which was a typical homofermentation to lactate (Figures 1, 4). How these fermentation profiles affect the feeding strategy of giant pandas and relate to its dietary shift to more pith during the spring time should be investigated more accurately with trials involving pandas, although in practice these are highly challenging. Such tests may consider *Bacteroides* (in green leaf) and *Escherichia/Shigella* (in yellow pith) as biomarkers indicative of solid lignocellulose degradation, as highlighted by the increase in their abundance upon removal of soluble organics (Supplementary Table S3).

When considering all metaproteins, the most abundant enzyme was the alpha amylase (E.C. 3.2.1.1) from the giant panda itself (Figure 7 and Supplementary Table S6). In de-H$_2$O yellow fermentations, where the solid yellow pith was the only carbon source, alpha-amylases constituted up to 60% of all the identified metaproteins in the sample, however, high levels were found in all tested condition (Figure 7). The copy number of alpha-amylase genes is higher in giant panda
Figure 7 | Relative amount of carbohydrate active enzymes (Cazy) at the end of laboratory-scale fermentation tests ($n = 3$) using microbiomes derived from giant panda fecal samples (Figure 2). E.C. numbers and relative abundances for Cazy reported in Supplementary Table S6; list of all metaproteins in Supplementary Table S7. Blue circles (bottom left) indicate the scale of relative abundance of all metaproteins (from 0.01 to 100%). Keys reported in the graph.

To summarize, the digestion of different portions of bamboo resulted in distinctive fermentation profiles. The green leaf was typically heterofermented to ethanol, lactate and acetate, while the yellow pith was homofermented to lactate. Microbial metabolic pathways inferred by metaproteomics suggested that hemicellulose, rather than cellulose or lignin, was the main source of energy. Profiling of fermentation products in stools and lab-scale cultivation of gut microbiomes to assess their actual biocatalytic potential should be implemented in large scale studies to explain the peculiar dietary behavior of giant pandas.

Materials and Methods

Inoculum Collection, Preparation, and Cultivation System

The fecal material was collected at the Pairi Daiza zoo (Brugellet, Belgium) and derived from the 6-year old male Xing Hui. No contact with the animal occurred during feces or bamboo collection. Entirely green or yellow stools were collected separately in 1 L airtight containers comprising an AnaeroGen™ bag (Oxoid, Hampshire, United Kingdom) to keep anoxic conditions until processing. About 5 kg of stools were collected.
for each color. Fresh *P. bisettii* bamboo as offered to Xing Hui was collected the same day (about 5 kg of fresh bamboo).

Stools were processed in the laboratory within 2 h following collection. For each stool, the external layer was removed. Stools containing a mixed color or traces of carrots or apples were discarded and only stools made entirely of either green leaves or yellow pith were selected. These were placed in autoclaved, anaerobic, milliQ water and stirred vigorously for 30 min while N\(_2\) was sparged to maintain anaerobic conditions. These inocula were termed ‘full inocula,’ as they contained microbes from the giant panda’s gut along with the soluble organics derived from panda’s gut digestion. Part of the full inocula from either green or yellow stools were used to generate a second inoculum termed ‘dewatered’ (de-H\(_2\)O) as follows: full inocula were placed in sterile falcon tubes and centrifuged for 10 min at 14000 rpm (Sorval RC5c PLUS, Beckman, Suarlée, Belgium); the supernatant was discarded and pellets resuspended in an equal volume of phosphate buffer saline solution (PBS), which had a pH of 7.35. The preparation method is summarized in Figure 2.

Green inocula were provided with green leaves of *P. bisettii*, while yellow inocula with yellow pith. The feeding strategy of Xing Hui was simulated when selecting leaves and pith. Leaves attached to thin, apical branches were collected. Leaves were milled into particles of about 1 mm using an electric grinder. The pith was generated by using a file on the yellow portion of the bamboo, which results from peeling off the green external cuticle of stems. The pith was also ground into particles of about 1 mm. Inocula and bamboo were incubated during 40 h in batch in serum bottles of 120 mL, with 50 mL of either green or yellow inocula provided with 1 g of green leaves or yellow pith, respectively. Reactors were capped using rubber stoppers and sealed with aluminum caps, their headspace was flushed with N\(_2\) for 15 min to keep anaerobiosis, and they were finally incubated at 37°C (the panda’s body temperature) in a shaking water bath (90 rpm, GLS Aqua 18 Plus, Grant).

**Microbiological Analysis**

Cell count and intact/damaged cell count was performed by flow cytometry. SYBR\(^{\text{R}}\) Green 1 (10\(^4\) concentrate in DMSO, Invitrogen) and Propidium Iodide (20 mM in dimethyl sulfoxide [DMSO], LIVE/DEAD BacLight Kit, Invitrogen, Belgium) were used to discriminate cells with intact and damaged cytoplasmic membranes. Flow cytometry was performed using a CyAn\(^{\text{TM}}\) ADP LX flow cytometer (Dakocytomation, Herlev, Belgium). MilliQ water was used as the sheath fluid. Data for 20000 events for each sample run was collected.

**Molecular Analysis**

DNA extraction for microbial community analysis was conducted on 1 g bamboo material, 1 g of fecal sample, or on the pellet resulting from 2 mL of either full inocula or de-H\(_2\)O reactors. The DNA was extracted with phenol-chloroform and precipitated with ice-cold isopropyl alcohol and 3M sodium acetate. DNA pellets were dried and resuspended in TE buffer and stored at −20°C. DNA quality was assessed using 1% (w/v) agarose gel electrophoresis (Life Technologies\(^{\text{TM}},\) ES), and quantified by a fluorescence assay (QuantiFluor\(^{\text{R}}\) dsDNA kit; Promega, United States) using a Glomax\(-\)Multi + system (Promega). Samples were normalized to 1 ng DNA µL\(^{-1}\) and sent to LGC Genomics (DE) for library preparation and sequencing via an Illumina Miseq platform (see Supplementary Information).

**Metaproteomics**

Culture samples (30 mL) were centrifuged, pellets resuspended in 2 mL 50 mM Tris/HCl (pH 6.8) and protein extracted with liquid phenol (Heyer et al., 2013). After protein quantification with amidol black assay, 25 µg of proteins were loaded into a 12% SDS-PAGE. The SDS-PAGE was conducted after proteins entered approximately 5 mm into the separation gel. The complete protein fraction was digested with trypsin, and peptides were measured by LC-MS/MS using an Elite Hybrid Ion Trap Orbitrap MS with a 120 min gradient. For protein identification, a database search with Mascot (Perkins et al., 1999) was performed, using a false discovery rate of 1% (see Supplementary Information).

**Chemical Analysis**

Lignocellulose content in bamboo was measured following the Van Soest method (Van Soest, 1963); TS and VS according Standard Method 2540G (APHA, 1992); and the pH using a pH probe (Herisau, Metrohm, Switzerland). Gas composition was analyzed with a Compact GC (Global Analyser Solutions, Breda, Netherlands), equipped with a Molsieve 5Å pre-column and two channels. Lactic and formic acid were determined with a 930 Compact IC Flex (Metrohm, Switzerland) ion chromatography (IC) system with inline bicarbonate removal (MCS), equipped with a guard column cartridge (Metrosep Dual 4/4.6, Metrhom) and an organic acid column (Metrosep 250/7.8, Metrohm) with a 850 IC conductivity detector. Alcoholics including glycerol and ethanol were determined with the same IC equipped with a guard column cartridge (Metrosep Trap 1 100/4.0, Metromh) and a Metrosep Carb 2 250/4.0 column (Metrohm) with an IC amperometric detector (see Supplementary Information).

**Statistical Analysis**

All statistical analyses were performed in the R statistical environment (v3.5.1) (R Core Team, 2015), using functions from the phyloseq (v1.16.2), DESeq2 (v1.22.1) and Phenolflow (v1.1) packages (McMurdie and Holmes, 2013; Love et al., 2014). Alpha diversity was assessed by the Hill diversity numbers, which incorporate both richness and evenness components (Hill, 1973). For beta diversity analysis the taxon abundances were rescaled by calculating their proportions and multiplying them by the minimum sample size present in the data set (McMurdie and Holmes, 2014). Beta diversity was then assessed by principal coordinate analysis (PCoA) of the Bray-Curtis dissimilarity matrix (see Supplementary Information). Results are the mean value of experiments made in 3–4 independent replicates, with error bars indicating standard deviation. Statistical significance was assessed using a non-parametric test (Mann–Whitney test) considering a two-sided distribution with 95% confidence interval.
DATA AVAILABILITY STATEMENT

The mass spectrometry proteomics data was deposited to the ProteomeXchange Consortium via the PRIDE partner repository with the dataset identifier PXD010872. The data on 16S sequences was deposited to NCBI under the BioProject ID: PRJNA574018.

ETHICS STATEMENT

Samples were collected under the strict supervision of Pairi Daiza zoo personnel. However, no contact with the animal occurred during feces or bamboo collection.

AUTHOR CONTRIBUTIONS

AS conceived, designed and performed the experiments, and wrote the manuscript. WK performed the experiments. MC designed and performed the experiments, and co-wrote the manuscript. RH performed the metaproteome analyses and co-wrote the manuscript. RP performed the 16S rRNA analyses. DB performed the metaproteome data analysis and co-wrote the manuscript. TB, DL, and HZ provided access to the giant panda fecal samples and bamboo. KR funded and supervised the project. All authors reviewed the manuscript.

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The remaining 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.

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