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Breath volatile metabolome reveals the impact of dietary fibres on the gut microbiota: Proof of concept in healthy volunteers

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Summary

Background Current data suggest that dietary fibre (DF) interaction with the gut microbiota largely contributes to their physiological effects. The bacterial fermentation of DF leads to the production of metabolites, most of them are volatile. This study analyzed the breath volatile metabolites (BVM) profile in healthy individuals (n=15) prior and after a 3-week intervention with chitin-glucan (CG, 4.5 g/day), an insoluble fermentable DF.

Methods The present exploratory study presents the original data related to the secondary outcomes, notably the analysis of BVM. BVM were analyzed throughout the test days -in fasting state and after standardized meals - using selected ion flow tube mass spectrometry (SIFT-MS). BVM production was correlated to the gut microbiota composition (Illumina sequencing, primary outcome), analyzed before and after the intervention.

Findings The data reveal that the post-prandial state versus fasting state is a key determinant of BVM fingerprint. Correlation analyses with fecal microbiota spotlighted butyrate-producing bacteria, notably Faecalibacterium, as dominant bacteria involved in butyrate and other BVM expiration. CG intervention promotes interindividual variations of fasting BVM, and decreases or delays the expiration of most exhaled BVM in favor of H₂ expiration, without any consequence on gastrointestinal tolerance.

Interpretation Assessing BVM is a non-invasive methodology allowing to analyze the influence of DF intervention on the gut microbiota.

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Keywords: Gut microbiota; Fibre; Short-chain fatty acids; Chitin-glucan; Fermentation; Breath volatile metabolome

Introduction The gut microbiota composition can be highly variable between individuals and the dietary habits explain around 20% of the microbial structural variations in humans. This highlights the potential for dietary strategies in disease management through gut microbiota
Research in context

Evidence before this study

The health effect of dietary fibres (DF) is of utmost importance considering the high capacity of the gut bacteria to produce bioactive metabolites upon DF fermentation. The identification of new biomarkers of DF intake through non-invasive analysis remains challenging. Breath analysis is a real-time, non-invasive and innovative method for detecting a wide range of volatile metabolites in expired breath. It is proposed as useful in disease diagnosis and as an indicator of metabolic disorders. However, the breath signature reflecting the interaction between DF intake and the gut microbiota has never been investigated until now.

Added value of this study

In the present exploratory study performed in healthy volunteers, we show that the nutritional status of the subjects is determinant to record the breath fingerprints, evaluated through selected ion flow tube mass spectrometry (SIFT-MS). The analysis reveals that several breath volatile metabolites (BVM) such as butyric acid, caproic acid, butanol, ethanol, phenol, methane, pentane, triethylamine, 3-hydroxybutanone and 2,3 butanedione can be exhaled in response to a standardized breakfast rich in DF. Most of those BVM are correlated with butyrate-producing bacteria, notably Faecalibacterium, a dominant gut bacteria, and may thus be proposed as biomarkers of the post-prandial gut bacterial metabolism. This study also shows that gut microbiota and exhaled BVM profile were modified after a chronic intake of chitin-glucan (CG), an insoluble DF whereas H2 expiration increased.

Implications of all the available evidence

Our results validate the hypothesis that the non-invasive breath analysis is an interesting tool to assess DF intake - gut microbiota interaction. It can be applicable for the follow up of most intervention studies targeting the gut microbiota for the improvement of health.

methods

Participants and study design

The FiberTAG study is an interventional monocentric study including two test days aiming to characterize gut microbiota composition and activity after a daily supplementation of 4.5 g of chitin-glucan (Kiotransine® from KitoZyme, Belgium) during 3 weeks, by analyzing volatile metabolites released in the breath and gut microbiota in stool samples. It is a longitudinal protocol, where the data were compared after 21 days of treatment versus time 0 in the same volunteers. Healthy young subjects were chosen to investigate the relevance of supplementing CG in the context of a healthy lifestyle. Sample size was estimated based on the primary outcome and using the software PASS 14.0.7 and the paired mean power modulation. Knowing that diet and microbiome are the strongest determinants of the human serum metabolism, the identification of bioactive metabolites issued from the interaction of food components and microbiota could be helpful in proposed biomarkers of health improvement based on nutritional strategies. Breath Volatile Metabolites (BVM) have been proposed as potential surrogate markers of gut dysbiosis in gastrointestinal diseases, as well as microbial signature of the fermentation of dietary fibre (DF).

Indeed, the gut bacteria are a major source of volatile metabolites such as methane, H2, short chain fatty acids (SCFA) or alkanes. Such metabolites are partly exhaled in breath. There are almost 900 volatile compounds in the exhaled breath of healthy humans. This suggests that BVM analysis could be an indicator of metabolic disorders, of specific dietary habits, or both. We have previously shown that a single intake of a DF (chitin glucan) supplementation in the morning, creates a specific BVM profile throughout the day in healthy volunteers.

The nutritional interest of DF comes from the recognition of their benefits for health based on an extensive literature. On the basis of studies in animals and humans, it has been proposed that DF might increase satiety, improve metabolic disorders, and modulate gut-related immunity through mechanisms related to SCFA production, thereby influencing endocrine and metabolic functions and intestinal epithelial integrity. Most DF are fermentable, entirely or at some extend and with different kinetics.

In the context of the FiberTAG project (Joint Programming Initiative “A Healthy Diet for a Healthy Life” 2017-2020 https://www.fibertag.eu/), we aimed at establishing a set of bioactive bacterial metabolites linking DF intake and gut microbiota-related health effect through non-invasive and innovative methods (therefore excluding blood sampling). A monocentric longitudinal intervention study intervention has been performed in healthy volunteers consisting in the daily administration of 4.5 g of chitin-glucan (CG), a novel insoluble DF considered as safe food ingredient by the European Food Safety Authority, for 3 weeks. We have demonstrated that after 3 weeks, such a supplementation with CG induces specific changes in the gut microbiota composition, promoting mainly Eubacterium, Dorea and Roseburia genera, and increased butyric, iso-valeric and caproic acids level in the faeces. Following those data, we hypothesized that BVM measurement after 3 weeks-CG intervention could reflect the interaction between DF intake and the gut microbiota. In the present article, we present data of BVM measurement performed by using selected ion flow tube mass spectrometry (SIFT-MS), in fasting state and in response to meals, the data being correlated to the fecal microbiome sequencing.
analysis test. Fifteen subjects are sufficient to detect with 94% of power a mean difference of 2% with an estimated standard deviation of difference of 2% (two-sided) in specific gut bacteria level (primary outcome). Of note, this number is in line with other studies focusing on breath metabolites analyses.19–21

Healthy subjects were recruited by the Center of Investigation in Clinical Nutrition (CICN-UCLouvain) by displaying posters on the university site and by mails, by social networks, local newspapers and local flyers disposed in shops and doctor’s offices in nearby cities. Thirty-four healthy males and females were recruited and underwent the screening test. Among them, sixteen subjects were included in the study based on the predefined inclusion and exclusion criteria and therefore received the allocated intervention. Fifteen completed the study.

Subjects were pre-screened by phone or mail using a questionnaire to test if they met the inclusion criteria (men or women, aged 18–40 years, body mass index (BMI) between 18 and 25 kg/m², Caucasian, non-smoker, in good general health as evidenced by medical history and physical examination, the use of effective contraception and physical examination, the use of effective contraception for women, and need to be H₂-producer. The exclusion criteria were: gastro-intestinal disorders, chronic or intestinal diseases (such as ulcers, diverticulitis or inflammatory bowel diseases), smoking, current or recent use of antibiotics, pro/prebiotics (as a dietary supplement) or any products affecting the gut transit (such as laxatives) within 4 weeks before starting the study, use of drugs that modify the composition of gut microbiota (antidiabetic drugs, cholesterol-lowering drugs, and proton pump inhibitors), pregnant or lactating woman (and woman who did not use highly effective contraception), food intolerance (e.g., lactose or gluten), presence of allergy or intolerance to one component of the tested product, excessive alcohol consumption (more than 3 units/day) and participation in another clinical trial 1 month before the screening visit. The recruitment was conducted from February to March 2018. The study was conducted from April to May 2018.

Thirty-four healthy males and females were recruited and underwent the screening test using lactulose to select H₂-producers as previously described and to exclude subjects with small intestinal bacteria overgrowth (SIBO).9 They were then invited to perform a screening test 2 weeks before the intervention. Food products consumption containing high amounts of dietary fibers (including whole grains, artichokes, Jerusalem artichokes, salsifies, leeks, onions) was limited to once a week. Moreover, subjects were asked to complete an electronic questionnaire (via a software called REDCap, Research Electronic Data Capture) every day during the intervention, which allowed monitoring subjects’ side effects and compliance. Empty and unused packets were returned to measure compliance.

Ethics

This study was approved by the local ethical committee (Comité d’Ethique Hospitalo-Facultaire UCLouvain/Cliniques Universitaires Saint-Luc; reference 2017/11SEP/437) and written informed consent was obtained from all subjects before inclusion in the study. The trial was carried out in accordance with the Good Clinical Practice (GCP) as required by the following regulations: the Belgian law of 7 May 2004 regarding experiments on the human persons and the EU Directive 2001/20/EC on Clinical Trials. The trial was registered at ClinicalTrials.gov under identification number NCT03505177; Registered 23 April 2018).

Study outcomes

The primary outcome of the study was the evaluation of 3-weeks of CG supplementation on the gut microbiota composition and results were already published by our team.28 The present exploratory study presents the
original data related to the secondary outcomes, notably the analysis of BVM upon a test day with highly controlled meal intake. Those data were obtained from the same subjects that were considered for the study of the primary outcome and previously presented.18

Test days
Both test days (day 0 and day 21) were organized in the same way. An overview of the study design is shown in Figure 1.

The evening before the two intervention days, subjects were asked to eat rice and meat, without vegetables and to avoid alcohol. Fasted subjects (from 9 pm the day before) arrived at 7 am at the investigation center (CICN). The study coordinator reviewed the inclusion and exclusion criteria and verified if instructions were adequately followed. After mouth wash (with Perio Aid Intensive Care), breath samples were taken to analyze BVM at baseline. Then, subject received a standardized breakfast rich in fibres (15 g in total) composed of 2 slices of rye bread coated with 25 g carob spread, and 200 ml orange juice. The breakfast has been eaten by all the participants within 15 min. Breath samples were collected at different times during 10 h after the breakfast ingestion to measure BVM. In the meantime, every hour, subjects were asked to a 100-mm VAS questionnaire at baseline and every hour for 10 h after the ingestion of the breakfast to investigate the 8 gastrointestinal symptoms (cramps, bloating, rumbling, discomfort, flatulence, burps, gastrointestinal reflux and nausea) usually described after the consumption of fibre. The scales were scored by measuring the distance (in mm) from 0 with a ruler. Scores were expressed as changes from baseline in cm. Of note, 2 h after the breakfast ingestion, subjects were free to drink water. Five hours after the start, the subjects received a standardized lunch composed of white bread, ham and cheese. The lunch has been eaten by all the participants within 15 min. At 4 pm (8 h after the start) subjects received a collation (sweet waffle). The test day ended at 6 pm.

Breath volatile metabolites (BVM) analysis
Breath samples (200 mL) were taken using specific sampling bags (FMONP.SBAG1.01 from MEC R&D SPRL, Belgium). The subjects were trained to blow deeply and slowly through the sampler after normal breathing. The bags were directly transported to Inter-science (Louvain-La-Neuve, Belgium) and then connected first to the SIFT-MS (Voice 200ultra, Syft Technologies, New Zealand) for BVM measurement, then to the GC using a TCD detector (Trace 1310 GC, Thermo Scientific, USA) for H2 analysis; all analyses being performed within 2 h after the harvest of the exhaled air as previously described.9 Three data sets of MS-fragments were obtained from H3O+, NO+ and O2+ ionizations (untargeted analysis). In addition, we targeted volatile metabolites known to study energy metabolism and to study the gut-related metabolic effects of DF, in particular those known to be a reflect of gut fermentation (coming from bacterial metabolism).5,21 Concentrations in ppm, determined by using an equation as previously described.9 Of note, we did not perform any data normalization step before identification of targeted BVM analyzed in the present study. Indeed, it was shown that several normalization approaches including probabilistic quotient normalization (PQN), were prone to introducing artefactual differences, in particular false positive biomarker identification and was not recommended for biomarkers identification through SIFT-MS technology.22,23 However, although it is known that the bags can have some inter-variations, we have considered BVM concentrations determined in bags filled with an inert gaz at each time point of the study, i.e. 0h-30min-2h-4h-5h-6h-8h-10h for both test days in order to remove a background bias that can appear due to environment and atmospheric conditions (such as temperature). Therefore, identified BVM (including H2) in breath samples were first corrected with the values obtained at each time point from those bags filled with helium (background removal). Then, we expressed data as absolute values in ppm or as changes from baseline in ppm.

Figure 1. Overview of the study design during the test days.
Data integration and statistical analysis
A per protocol analysis was completed for this study. Data are expressed as medians ± interquartile ranges. However, since the median values are 0 for most of the gastrointestinal symptoms whatever the time considered, data are expressed also as means ± SD for Table S3 and Figure S5. Net area under the curve (AUC) were calculated for each symptom recorded every hour and for each identified BVM recorded during 10 h. Wilcoxon matched-pairs signed rank test were used to compare net AUC. In addition, mixed model followed by Sidak post-test were performed to compare evolution BVM over time (Figure S2). PCA model (presented in the Figure S1 and the supplementary video) was built on raw data without any normalization step by using R (v3.5.1, package “ade4”). Correlation analysis was performed between BVM and genera, with a mean of relative abundance ≥ 1% at day 0 or day 21, by Spearman’s correlation test using the “corrplot” package in R. A significance level of \( p < 0.05 \) was adopted for all analyses.

Role of the funding source
The funding source had no role in study design, data collection, data analysis, data interpretation, writing of the report and in the decision to submit the paper for publication. The corresponding author confirms that she had full access to all the data of the study and had final responsibility for the decision to submit for publication.

Results
Subjects
Seven men and eight women were selected as H\(_2\)-producers based on the lactulose test (data not shown) and completed the study respecting inclusion and exclusion criteria; only one volunteer dropped out on the last test day (no reason was given). There was 96.5% compliance in the participants. Baseline characteristics of participants are shown in Table S1. Carbohydrate, lipid, protein, fibre and alcohol intakes were not significantly affected whereas energy intake decreased by approximately 15% after 3 weeks of CG supplementation.

Higher H\(_2\) was expired in response to the ingestion of a breakfast rich in DF after CG treatment
The Figure 2 presents H\(_2\) exhaled in the breath and measured with GC-TCD at time 0h, 5h, and 10h. H\(_2\) was exhaled with concentrations higher than 200 ppm. Basal levels obtained at fasting state (time 0h) were significantly different between both test days (\( p = 0.0186 \)). Interestingly, exhaled H\(_2\) decreased 5h and 10h (-20 ppm) after the breakfast intake at day 0 whereas it increased (+ 20 ppm) after CG intervention (\( p = 0.0001 \)).

BVM profile in the breath was modified in response to the ingestion of a breakfast rich in DF after CG treatment
Untargeted analysis using SIFT-MS was performed to record the breath fingerprints at different times during the test days. The dynamic evolution of MS-fragments obtained by ionisation with three precursor ions (H\(_3\)O\(^+\), NO\(^+\), and O\(_2\)\(^+\)) was shown through principal component analysis (PCA) analysis (Figure S1 and supplementary video). We observed that separate clusters were formed depending on the time of measurement for the 15 subjects at both test days. The most discriminant time was the overnight-fasted state (time 0h) versus fed state (from time 30 min), pre-prandial state (meaning before lunch, time 4h and 5h) or post-prandial-post absorptive state (from time 6h). Furthermore, we observed that the individual variability was greater at
overnight-fasted state, an effect promoted after CG treatment during 3 weeks.

A targeted analysis was conducted to identify 24 BVM, namely issued from bacterial metabolism. The evolution of all BVM exhaled in the breath over time prior and after 3 weeks of CG intake are presented in Figure S2 in absolute values (ppm) whereas Figures 3 and S3 presented the concentrations in ppm corrected by the baseline values obtained at time oh. We observed a large variation in concentration from one BVM exhaled in the breath to another from 0.001 to 100 ppm (Figure S3). We found that only two BVM -methane and acetone- were detected at levels higher than 1 ppm and both were not affected by CG intake. Among BVM detected in the breath at lower levels (between 0.01 and 1 ppm), acetaldehyde and phenol expired at fasted state (time 0h) were significantly higher at day 21 versus day 0 whereas fasting exhaled pentane, butyric acid and 3-hydroxybutanone were lower after the intervention (mixed model: $p < 0.0001$ for treatment). Exhaled H$_2$S, triethylamine and caproic acid detected with levels below 0.01 ppm were also different at time 0h between both test days.

We focused our study on the evolution of exhaled BVM produced -expressed as change versus the baseline value in fasted state- after ingestion of a breakfast rich in

![Figure 3. Targeted BVM concentrations (changes from baseline) exhaled in breath of healthy subjects (n=15) in response to a standardized breakfast intake prior (day 0) and after (day 21) chitin-glucan intake (and significantly affected by the chitin-glucan intake during 3 weeks). Data are medians ± interquartile ranges (*p < 0.05; matched-pairs Wilcoxon signed-rank test on net area under the curve).](image-url)
DF (Figures 3 and S3). Mixed model revealed a significant effect of time for all targeted BVM ($p < 0.05$). A significant effect of CG treatment was obtained for butyric acid, caproic acid, ethanol, propionoic acid, methane, pentane, triethylamine, 2,3-butanedione, and acetaldehyde (mixed model performed on all BVM evolution: $p < 0.05$). Among 6 SCFA identified, the net AUC of the evolution of butyric and caproic acid concentrations after the DF-rich breakfast was significantly decreased by the CG treatment ($p = 0.0353$ and $p = 0.0004$, respectively) (Figure 3). Besides changes in SCFA exhaled in the breath, Figure 3 revealed 8 other BVM with a net AUC significantly lower after the nutritional intervention with CG, that are butanol ($p = 0.0084$), ethanol ($p = 0.0034$), phenol ($p < 0.0001$), methane ($p = 0.0151$), pentane ($p = 0.0001$), triethylamine ($p = 0.0215$), 2,3-butanedione ($p < 0.0001$) and acetaldehyde ($p = 0.0003$). The power calculation and the effect size on those secondary outcomes were presented in the Table S2. We observed that the power and the size effect are excellent ($>80\%$ and $>0.8$, respectively) for 6 of them. Propionic acid and acetaldehyde were the sole BVM that decreased in response to the breakfast without any other variation observed throughout the day. The decrease of acetaldehyde was more pronounced after CG treatment whereas the level of exhaled propionic acid remained higher all day long after CG intake.

Overall, it is important to note that for some BVM, their level was higher in the fasting than in all measurement performed later on, in the post-prandial or post-absorptive state. It is the case for propionic acid, propanol and benzene that drastically dropped (by more than 40-fold for propionic acid, and around 10-fold for propanol) after the breakfast (Figures 3, S3). Acetic acid and acetaldehyde, also dropped down at a lesser extend (by about 2-fold) upon feeding. All the other BVM remained stable or increased at different time points during the test days.

Concerning exhaled SCFA, butyric, caproic and (iso)valeric acids increased just after the breakfast (mostly at time 2h) before CG intervention with a return to basal values only for (iso)valeric acids, whereas butyric and caproic acids continued to rise (Figures 3, S3). Furthermore, the highest level of SCFA exhaled in the breath (except for acetic and propionic acids that decreased) were obtained much later, i.e. at least 8h after the standard breakfast when CG was consumed during weeks.

As observed for butyric acid, caproic acid, butanol, ethanol, phenol, methane, pentane, triethylamine, pentane, and 2,3-butanedione increased in response to fibre-rich breakfast at day 0 before CG intervention, with the highest levels obtained mainly 8h after breakfast intake (Figure 3). Surprisingly, CG consumption during 3 weeks reduced the magnitude of the response to the DF-enriched breakfast for those microbial metabolites at 8h ($^*p < 0.05$; matched-pairs Wilcoxon signed-rank test performed at time 8h).

Although similar evolution of BVM were observed for other targeted BVM (e.g. hydroxybutanone followed the same trend than butyric acid), matched-pairs Wilcoxon signed-rank test performed on total net AUC did not reveal any significant effect of CG intervention (Figure S3).

Importantly, BVM were differently affected considering the post-prandial post-breakfast or post-lunch states (Figure S4). Most significant changes occurred in the post-lunch state upon the CG supplementation: among BVM that increased after the lunch versus the fasted state for both days, higher levels of 2-methylbutyric acid, isovaleric acid, acetone exhaled in the breath whereas lower levels of triethylamine, caproic acid, butyric acid, butanol, ethanol, phenol, pentane and methane were observed after CG treatment (Figure S4b).

Gastrointestinal tolerance after ingestion of a breakfast rich in DF was not modified after the CG treatment

Knowing that hydrogen production after feeding was higher after CG treatment, and that this DF is largely fermented, the question of the potential effect of CG treatment on gastrointestinal symptoms merited to be evaluated. We show that the baseline scores of gastrointestinal parameters before and after CG treatment were not significantly different ($p > 0.05$) (Table S3). Gastrointestinal tolerance was investigated after ingesting of the standardised breakfast rich in fibre. The evolution of the different scores throughout the day was not significantly affected by CG intake (matched-pairs Wilcoxon signed-rank test on net AUC: $p > 0.05$) (Figure S5).

Correlation analyses between exhaled BVM and gut microbiota composition

The gut microbiota of participants was analyzed before and after 3 weeks of CG intervention. \(^{18}\) Figure 4a shows the relative abundance at the genus level at baseline prior any DF intervention. We observed that 50% of the fecal bacteria of healthy subjects were dominated by 9 genera, most of them being butyrate-producing bacteria (Faecalibacterium, Ruminococcus, Eubacterium, unclassified Lachnospiraceae, Coprococcus). Correlation analysis was performed between gut microbiota composition and BVM prior CG intervention (day 0) considering the time where the highest levels of BVM were obtained, i.e 8h after the breakfast. Interestingly, Faecalibacterium, the most abundant one (7 % in relative abundance) was correlated ($p = 0.0498$) with exhaled butyric acid 8h after the breakfast intake (Figure 4b). Moreover, this genus was also positively correlated with other BVM, most of them being bacterial metabolites: butanol ($p = 0.0351$), acetaldehyde ($p = 0.0260$), ethanol ($p = 0.0351$), isoprene ($p = 0.0019$), propionic acid ($0.0189$), caproic acid ($0.0498$) and 3-hydroxybutanone...
At that time, other exhaled bacterial metabolites (propionic acid, 2,3 butanedione, phenol, pentane, methane) were positively correlated with abundant genera (>1%), mainly Prevotella or genera from Ruminococcaceae and Lachnospiraceae families (p < 0.05).

The impact of CG intake on gut microbial changes after 3 weeks of intervention was previously described.18 We concluded that a 3-week CG supplementation did not induce major changes in the overall gut microbiota composition (no change in α-diversity and β-diversity indexes). We have previously shown that CG consumption for 3 weeks resulted in specific changes to bacterial genera and species.18 Indeed, we found that 3 weeks of CG intake significantly increased Lachnospiraceae UCG-004, Roseburia, Eubacterium, Ruminococcaceae UCG-003 and UCG-005, whereas it decreased Subdoligranulum, Bergeyella, Blautia and Dorea. However, among bacteria that increased due to CG intervention during 3 weeks (and accounting for more than 1% for each subject), none of those were positively related to expiration of bacterial metabolites (see correlation analyses between BVM at time 8h and genera at D21 in Supplementary data).

Discussion

In general, metabolites measured in biological fluids are considered as interesting biomarkers of human clinical and nutritional status and can therefore be proposed as a useful diagnostic aid for monitoring metabolic diseases.6 Breath analysis is a non-invasive and sensitive medical diagnostic tool that offers the possibility of rapid measurements.8 For example, BVM analysis hence serve as attractive, non-invasive biomarkers for diagnosing and monitoring irritable bowel syndrome and inflammatory bowel disease.24 However, the origin of volatile metabolites is not yet fully understood. Analytical methods are continually developed to allow for better detection and quantification of volatile metabolites, thus enabling the detection of a wider range of biomarkers of pathophysiology and nutritional status.
Direct MS techniques such as SIFT-MS have gained increasing interest because they offer the promise of selectivity, speed and sensitivity (low parts per billion by volume levels), and SIFT-MS is particularly well suited for the purpose of exhaled breath analysis. This technique can be used in untargeted scan mode, where it provides pattern-based classification capacity but usually, it is applied in targeted mode, which means that only a limited set of components are measured simultaneously. The contribution of nutrition in the profile of BVM remains poorly evaluated. Some studies monitored exhaled breath upon changes in dietary habits (high \textit{versus} low fat dairy drink, high \textit{versus} low fibre diets, gluten-free diet). Recently, we have analysed the impact of a single intake of an insoluble DF (4.5 g CG) on breath volatile metabolome in healthy subjects. The targeted approach obtained with SIFT-MS analysis and correlation analyses with gut microbiota composition suggested that exhaled butyric acid, triethylamine, pentane, 3-hydroxybutanone and 2,3-butanedione were identified as microbial signature of CG intake with the highest levels observed 6h after its ingestion. In a recent study, it was shown that no significant changes in volatile compound profiles (using thermal desorption gas chromatography coupled to time-of-flight mass spectrometry) were found after 4 weeks of a prebiotic dietary fibre supplementation (galacto-oligosaccharides, 15g/day) both in elderly as well as in young adults after an overnight fast, indicating relatively small differences in exhaled breath profiles, which is in line with microbiota composition and fecal metabolite data. The objective of the present exploratory study was to investigate the complex interaction between DF and gut microbiota by assessing BVM before and after 3 weeks of CG consumption in a representative homogeneous population of healthy young gender-mixed volunteers with the same level of education. The recruitment of subjects performed on a university site allowed to reach this objective since 7 men and 8 women with a mean age of 21 in good general health, as evidenced by medical history and physical examination, were included in the study. We have applied the SIFT-MS methodology in order to follow exhaled BVM in response to a standardized breakfast rich in fibre prior and after chronic consumption of CG (4.5g per day during 21 days). There was an excellent compliance of the participants reaching 96.5%. Thus, this parameter does not impact on study inferences. We selected rye bread and carob spread to design the test meal because of their DF richness and fermentation capacity was already demonstrated. Indeed, carob spread is a high DF and polyphenol containing product where caproic acid has been recently shown to be
produced from lactate oxidation.\textsuperscript{16,37} In bacteria, lactate oxidation provides acetyl-CoA which enters elongation process to form butyryl-CoA and hexanoyl-CoA (also called caproyl-CoA).

Besides SCFA, a rise in butanol, ethanol, phenol, methane, pentane, triethylamine, 2,3-butanedione, acetone, acetonitrile, benzaldehyde, 3-hydroxybutanone, H$_2$S or methanol also occurred after lunch before CG intervention (at day 0) and often peaked after 8 h. For these BVM, it seems relevant to propose the gut fermentation of nutrients reaching the colon as the driver of their production and release in the breath. Therefore, the correlation analyses were performed between BVM (in ppm, corrected by the baseline value at fasted state) at the time where most of breath BVM were present in the breath at the highest levels, i.e. 8h after the breakfast intake and genera (with relative abundance > 1%). Butyric and caproic acids were positively correlated to \textit{Faecalibacterium}, the most abundant genus detected in healthy subjects at baseline. This genus was also correlated with high levels (obtained at time 8h) of other exhaled BVM recognized as bacterial metabolites such as butanol, ethanol and 3-hydroxybutanone.\textsuperscript{38,39} Furthermore, 2.3 butanediol, phenol, pentane and methane are other bacterial metabolites exhaled in the breath and were positively correlated with the most interesting butyrate-producing bacteria, mainly from \textit{Ruminococcaceae} and \textit{Lachnospiraceae} families. Bacterial production of 3-hydroxybutanone and 2.3-butanediol is related to citrate metabolism through the citrate—oxaloacetate—pyruvate—acetolactate—acetoin/diacetyl pathway, where pyruvate is considered as the precursor.\textsuperscript{39} Alcohol/aldehyde dehydrogenase found in some \textit{Clostridia} converts butyryl-CoA to n-butanol.\textsuperscript{39} While the origin of breath methane is undoubtedly linked to gut fermentation, intestinal bacteria are also considered as a major source of breath pentane versus the contribution of endogenous membrane lipid peroxidation.\textsuperscript{5,4,42} It should be interesting to have data supporting the relative contribution of bacterial versus host metabolism in the production of the metabolites such as pentane. Indeed, it was already shown that exhaled pentane was dependent on dietary linoleate intake, and was blunted upon antibiotic treatment.\textsuperscript{8,42} Altogether, our data suggest that SCFA such as butyric and caproic acids but also other metabolites such as ethanol, butanol, phenol, 3-hydroxybutanone, 2,3 butanediol, pentane and methane exhaled in the breath are issued from bacterial metabolism in the lower part of the gut, depending mostly on bacteria known to be butyrate-producers. It should be interesting to know if the metabolic pathways present the identified bacteria are able to produce such metabolites. Undeniably, further studies are required to study the origin and the fate of most BVM metabolites.

We have demonstrated that the shift in the gut microbiota composition occurring upon CG intervention during 3 weeks (mainly increase of \textit{Roseburia} and \textit{Eubacterium} genera) is not due to intra-individual variations of microbiome with time and could clearly be attributed to CG treatment, making those bacteria privileged targets of CG intake.\textsuperscript{18} In fact, \textit{Roseburia} is known for their important ability to utilize diet-derived polysaccharides and in turn produce butyrate.\textsuperscript{39} Consistently, we highlighted higher fecal butyric acid concentration after CG supplementation.\textsuperscript{18} However, this effect was not accompanied by higher expiration of this SCFA in the breath in response to a meal rich in DF. We observed that C$_5$ and C$_6$ SCFA were also affected by the dietary intervention but at much lesser extent (delta < 0.01 ppm). Besides SCFA exhaled in the breath, 7 BVM reflecting bacterial metabolism in the colon were significantly affected by the nutritional intervention with CG. Indeed, butanol, ethanol, phenol, methane, pentane, triethylamine and 2,3-butandione were exhaled with highest levels obtained at least 6h after the DF-enriched breakfast but the amplitude of the response was much lower after the CG supplementation. Of particular interest, analysis focusing on post-lunch state revealed that exhaled butyric acid, caproic acid, butanol, ethanol, phenol, triethylamine, pentane and methane decreased after CG intervention. In contrast, breath H$_2$ increased after CG intervention in response to the DF-enriched breakfast. We observed that the basal level of H$_2$ release in the fasting state after the treatment were significantly different between both test days. Knowing that the evening meal – and its content in DF- may influence the H$_2$ production in the next morning,\textsuperscript{43} we have standardized the evening meal (that consisted in eating meat, rice without vegetables) in order to minimize DF intake. Therefore, the different levels in fasting state at day 0 and day 21, could be due to the modifications of the gut microbiota composition (and function) linked to chronic CG intake and the potential fermentation of nutrients in the large intestine during the night. In fact, microbial H$_2$ cycling is central to metabolic homeostasis and microbial composition in the human gastrointestinal tract. Molecular H$_2$ is produced as an end product of carbohydrate fermentation, and may be subsequently utilized by cross-feeding microbes for growth and in the production of larger molecules (H$_2$ is reoxidised primarily by sulfate-reduction, acetogenesis, and methanogenesis).\textsuperscript{42,44} Wolf et al. have demonstrated that 70% of gastrointestinal microbial species encode the genetic capacity to metabolise H$_2$ and its relative proportion compared to the other bacterial products depend on the environmental conditions (e.g. pH).\textsuperscript{44} Furthermore, the presence of H$_2$ also produces a thermodynamic environment favouring SCFA production.\textsuperscript{44} We have also to consider that more than 90% of the colon bacteria are butyrate-producers represented by \textit{Eubacterium/Roseburia} spp., which belong to clostridial cluster XIVa (24% of the total colon microbiota), and \textit{Faecalibacterium prausnitzii}, which belongs to the clostridial cluster IV (7% of the total colon microbiota).\textsuperscript{45} A quantitative link or stoichiometrically balanced pathway between bacterial substrate metabolism and H$_2$ gas formation have already been proposed.
in cluster XIVa butyrate-producing colon bacteria.\textsuperscript{45,46} Considering the higher proportion of two important butyrate-producers (\textit{Eubacterium} and \textit{Roseburia}) observed after chronic consumption of CG, those links were probably involved in mechanisms explaining the higher production of breath H\textsubscript{2}.

We have identified several limitations to our study. First, it is a longitudinal protocol, where the data were compared after 21 days of treatment \textit{versus} time 0 in the same subjects. This kind of strategy has already been used in order to highlight microbial signature of a probiotic DF.\textsuperscript{47} We cannot rule out that some changes in BVM may have occurred with time independent of the treatment. Secondly, this study is focused on a small cohort investigated in a single center. Thirdly, it potentially included some bias by self-reporting of food diaries. Finally, hydrogen-producers only (and excluding subjects with SIBO) were considered.

In conclusion, this study showed that several bacterial metabolites can be exhaled in the breath in response to a meal rich in DF. Although we cannot strictly exclude the presence of an altered metabolic homeostasis and/or an altered gut microbiota, it is important to note that medical history, drug history, inclusion and exclusion criteria and some clinical parameters obtained at the baseline led us to exclude metabolic diseases known to be link with gut dysbiosis such as obesity, diabetes or hypertension, and to suppose this young population as “healthy”. This exploratory study identified exhaled butyric acid, caproic acid, butanol, ethanol, phenol, methane, pentane, triethylamine, 3-hydroxybutanoic and 2,3 butanedione as biomarkers of the gut bacterial metabolism after a breakfast rich in DF. We have to note that the power is less than 80\% only for 4 out of those 10 BVM. Our data suggested that the chronic intake of CG, an insoluble DF, decreased or delayed the expiration of most exhaled BVM in favor of H\textsubscript{2} expiration in response to a meal rich in DF, without inducing uncomfortable gastrointestinal symptoms. The lower exhalation of BVM such as triethylamine, acetaldehyde and pentane could be interesting to explore in the context of inflammatory bowel disease (Crohn’s disease, ulcerative colitis) or nonalcoholic fatty liver disease (NAFLD) since their level can be found significantly elevated in the breath of those patients.\textsuperscript{48-49} Altogether with our previous study devoted to study breath volatile metabolome after a single intake of CG,\textsuperscript{9} we confirmed that analysis of BVM in exhaled air can be used as a dynamic approach to noninvasively assess the complex interaction between DF and gut microbiota. Our data also support the interest of focusing on the role of nutrition on exhaled BVM profiling at a time where this non-invasive procedure is proposed as an interesting tool to unravel metabolic and infectious diseases, like recently shown for Covid 19 infection in human.\textsuperscript{30} This study supports the relevance of breath volatile metabolomic approach to unravel nutrients-gut microbiota-host interactions. Further experiments with different types of DF would allow to unravel the metabolomic signature of their interaction with the gut microbiome, knowing that the concepts of selective fermentation of DF is largely studied \textit{in vitro}, but poorly assessed \textit{in vivo}. The non-invasive sampling required for this type of analysis also appears as a positive outcome of our data.

**Contributors**

Conceptualization: A.M.N., N.M.D.; Data curation: A.M.N., J.R.; Formal analysis: A.M.N., J.R., Z.Z.; Funding acquisition: N.M.D., V.M., M.L., S.C.B., J.W.; Investigation: A.M.N., J.R. Methodology: A.M.N., J.R., Z.Z., L.B. B.; Project administration: N.M.D., A.N.M.; Resources: P.D.C., N.M.D., V.M.; Software: J.R.; Supervision: N.M. D.; Validation with verification of the underlying data: N.M.D., A.M.N., J.R.; Visualization: A.M.N., J.R.; Writing—original draft: A.M.N.; Writing—review and editing: J.R., Z.Z., J-A.N., P.D.C., L.B.B., N.M.D. All authors read and approved the final version of the manuscript.

**Declaration of interests**

P.D.C. is cofounder of A-Mansia Biotech SA and owner of patents on Akkermansia muciniphila, gut microbes and metabolic diseases. The other authors declare no conflict of interest.

**Data sharing statement**

The study protocol and the datasets generated during and/or analyzed during the current study, including de-identified participant data will be available from the corresponding author on reasonable request.

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Supplementary materials

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