The Co-Regulation of the Gut Microbiome and Host Genes Might Play Important Roles in Metformin Intolerance

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Research

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Abstract

Background

Metformin is commonly considered the first-line therapy for type 2 diabetes (T2D) and also had potential treating utility in other areas; however, ~20% of patients experience intolerance with unclear underlying mechanisms. In the present study, we performed the full-length 16S rRNA (V1-V9) for the fecal samples and bioinformatics analysis to study the mechanisms of the metformin intolerance combining the gut flora and host.

Results

The results showed that *Barnesiella* (p=0.046) and *Parabacteroides goldsteinii* (p=0.016), which transforming primary into secondary bile acid (SBA), were higher in the TS than T group, and were eliminated in the TSa group, which might lead to the accumulation of primary bile acids (PBA) such as cholic acid (CA), the change of *GLI1* gene, and following diarrhea in the TSa group. *Lactobacillus brevis* (p=0.024) and *Lactobacillus plantarum* (p=0.026) were up-regulated in TSa than TS group. The two flora might cause the changes of genes including *FOXA2*, *HTR7*, *GADPH*, and intolerance relief, which might be a worthwhile future direction for preventing metformin intolerance.

Conclusions

These results hinted that the differential flora and co-regulation of them with the host might be intolerance-related. Our results partly provided theoretical support for intolerance prevention.

Background

Metformin is commonly considered as the first-line therapy for type 2 diabetes (T2D) and also had potential treating utility in obesity, metabolic dysfunction, and some cancers. However, ~20% of patients experience gastrointestinal (GI) intolerance, including diarrhea, nausea, and bloating with unclear underlying mechanisms and a lack of effective management strategies that warrant discontinuation of metformin treatment [1]. Researches have uncovered that the modification of intestinal microbiota and host DNA methylation might be both involved in GI intolerance, and the increase of *Escherichia* abundance might be one of the reasons in the European population. However, the *Escherichia* have not been modified in the Chinese population [2], which implied that different mechanisms might employed in Chinese.

In present study, we collected fecal samples from T2D patients who had no GI intolerance after metformin administration (before (T) and after (Ta) taking metformin, respectively) and who had GI intolerance after metformin administration (before (TS) and after (TSa) taking metformin, respectively)
and health subjects (N). Then, we carried out 16S rRNA sequencing on fecal samples and bioinformatics analysis to explore the mechanisms of metformin intolerance.

**Results And Discussion**

We found that the *Dorea longicatena* was up-regulated in both T and TS groups (LDA Score>2.0) and also in the merging data than N group (Figure S1a-c). The K03386, K01809, K03321, and K13016 are enriched in the T group, and K02398 is enriched in the N group (Table 1). The alpha diversity was not differed before vs. after taking metformin (Table S2). These patterns had been confirmed by previous studies [3, 4], which proved the high confidence of our research.

| Group  | KO Class                              | KO ID and Definition                                                                 |
|--------|--------------------------------------|--------------------------------------------------------------------------------------|
| T-N    | KOs up-regulated in T group          | K03386: PRDX2_4, ahpC; peroxiredoxin (alkyl hydroperoxide reductase subunit C) [EC:1.11.1.15] |
|        |                                      | K01809: manA, MPI; mannose-6-phosphate isomerase [EC:5.3.1.8]                        |
|        |                                      | K03321: TC.SULP; sulfate permease, SulP family                                       |
|        |                                      | K13016: wbpB; UDP-N-acetyl-2-amino-2-deoxyglucuronate dehydrogenase [EC:1.1.1.335] |
|        | KOs up-regulated in N group          | K02398: flgM; negative regulator of flagellin synthesis FlgM                         |
| T-TS   | Flagellin-related KOs up-regulated in TS groups | K02409: fliF; flagellar M-ring protein                                                |
|        |                                      | K02389: flgD; flagellar basal-body rod modification protein                          |
|        |                                      | K02397: flgL; flagellar hook-associated protein 3                                    |
|        |                                      | K02413: flij; flagellar FliJ protein                                                 |
|        |                                      | K02398: flgM; negative regulator of flagellin synthesis                                |
|        |                                      | K02395: flgJ; flagellar protein                                                     |
| TS-TSa | Lactose degradation related KO down-regulated in TSa than TS group                   | K01190: lacZ; beta-galactosidase [EC:3.2.1.23]                                        |
| T-Ta   | RFO degradation related KO up-regulated in Ta than T groups                          | K07406: melA; alpha-galactosidase [EC:3.2.1.22]                                       |

Importantly, we found that the background difference and changes caused by metformin in the gut microbiome might be both involved in GI intolerance. The results showed that the *Barnesiella* (p=0.046)
and Parabacteroides goldsteinii (p=0.016), which transforming primary into secondary bile acids (SBA) [5, 6] were far higher in the TS than T group. However, the two flora were eliminated in the TSa group (Figure 1a), which implied that the eradication of Barnesiella and P. goldsteinii might lead to the accumulation of PBA such as CA and following diarrhea in the TSa group (Figure 2 inside (1) dashed rectangle). Six flagella assembly-relevant Kyoto Encyclopedia of Genes and Genomes (KEGG) orthologs (KOs) significantly higher enriched in TS than T group were also found (Table 1), which might cause a higher risk of inflammation in the TS group for the pro-inflammatory potential of flagellin [7]. The TS and T groups exhibited different patterns of microbial communities from TSa and Ta groups (Figure 1b).

There are 26 and 10 significantly differential flora in total between the TS and TSa group, and the T and Ta group, respectively (Figure 1c, Table S2,3). Among them, Leuconostocaceae (p=0.048), Leuconostoc (p=0.041), Weissella confusa (p=0.037), Weissella paramesenteroides (p=0.008), Lactobacillus brevis (p=0.024), Lactobacillus plantarum (p=0.026) were up-regulated in TSa group (Figure 1d). These flora belongs to lactic acid bacteria. In addition, the alpha-galactosidase, a protective factor of GI intolerance [8], was higher enriched in Ta than T group but not in the TSa group. In contrast, beta-galactosidase, also a protective factor of GI intolerance [9], was down-regulated in TSa compared with the TS group (Table 1). It may be one reason for GI intolerance (Figure 2, represented by black and pink lines respectively inside (2) dashed ellipse).

Besides, by bioinformatics analysis as that showed in Supplementary, we found that metformin intolerance might be the result of co-regulation of intestinal flora and host genes. There is a close-interaction among the L. plantarum, L. brevis, cholic acid (CA) and, metformin glycemic response and intolerance-related genes (Figure 1e, Table S4-6). According to GEO data and previous reports, we found that the CA and FOXA2 could both modulate the expression of GLI1 [10]. In contrast, the L. plantarum and L. brevis could respectively modulate the expression of HTR7 and GAPDH. By RT-qPCR check, we found the expression of GLI1 was down-regulated (p<0.05), and the expression of FOXA2 showed an upward trend (p<0.10) in TSa than TS group (Figure 2, Figure S3, Table S7). The downregulation of GLI1 might cause bloating, diarrhea and nausea by sharing the same pathogenesis and symptoms of inflammatory bowel disease (IBD), which might lead to GI intolerance [11]. On the contrary, we found expression of HTR7 and GADPH showed a downward trend (p<0.10) in TSa than TS group (Figure 2, Figure S3, Table S7), which might be related to the relief of GI intolerance, for opposite changes of these genes are corresponding to IBD [12, 13]. The effect of flora on these genes seem to restore a healthy state after being disturbed, which might be one of the reasons why some intolerance individuals become more tolerant to metformin after persisting on it for a period, which might be a worthwhile future direction for the prevention of metformin GI intolerance.

**Conclusion**

In general, the mechanism of metformin intolerance in Chinese might be different from that in Europeans. It might be caused by the eradication of PBA degrading-related flora in gut microbiota and also the co-regulation of intestinal flora and host genes. However, the relatively small sample size might affect the results to a certain extent. However, fortunately, some of the results in our study are consistent with large
cohort researches, which guaranteeing the accuracy of the results. More importantly, the selection of dose regimen and treatment duration was based on the hospital's conventional treatment process to avoid interference treatment and better reflect the actual clinical phenomenon. Therefore, our results could still provide theoretical support for intolerance prevention. Further experiments with larger sample sizes and animal models are required to verify those results to reveal the mechanism involved so that more people could benefit from metformin in the future.

**Material And Methods**

**Subjects**

This study was carried out at the First People's Hospital of Qujing City, Yunnan Province, China, from February 2019 to August 2019. The inclusion criteria were as follows: the patients were newly diagnosed with T2D with the fasting plasma (blood) glucose higher than 7.0 mmol/L, were between 40 to 70 years old, were able to communicate, had volunteered to participate in this study, and were willing to provide informed consent. Subjects did not take ion pump inhibitor drugs, antibiotics, steroid hormones, or Chinese herbal medicine, including oral, intramuscular, or intravenous injections within the three months before collecting fecal samples, did not take other glucose-lowering medications. Subjects did not take other medicine except drugs used in this study during the experiment, did not occur diarrhea on the day of the first sampling. Those excluded were the patients who had severe conditions, including indigestion, renal failure, hepatic failure, severe gallbladder, stroke, pancreatic diseases, malignant tumors, or unstable cardiovascular diseases (such as myocardial infarction, ketosis, or hyperthyroidism) [14]. Age-matched healthy volunteers were included as above.

**Medication Strategy and Samples Collection**

We recommend the use of an oral glucose tolerance test (OGTT) (consisting of a fasting and 2-hour glucose level using a 75-g oral glucose load) to screen for impaired glucose tolerance (IGT) and T2D. Feces and blood samples were collected, and the OGTT experiment was completed in the early morning of the next day after the patient was admitted to the hospital. The blood was used to detect other indicators such as fasting blood glucose, serum C-peptide. Then the patient took metformin hydrochloride sustained-release tablet (Qingdao Huanghai Pharmaceutical Co., Ltd.) orally at a dose of 500 mg/time, two times/day. When the patient had intestinal side effects, they stopped metformin treatment, collected stool, and measured fasting blood glucose the next morning. When the patient had no side effects after metformin administration, the stool was collected five days later, and fasting blood glucose was measured. The feces of each subject were immediately stored at -80°C after collection until the next step. According to hospital clinical experience, insulin combined with metformin treatment can achieve a better hypoglycemic effect, and insulin will not change the composition of intestinal flora [2]. So the patient had been treated with an insulin pump with the weight (kg) * 0.2-0.5 u/day dose first for a day, when whose random blood glucose was greater than 16.8 mmol/L on admission. Feces and blood samples were collected from the healthy subjects only one time, respectively.
Isolation and qualification of fecal bacterial DNA

Eighteen stool samples were collected from twelve subjects (Table S1), as follows: six samples from three patients who had no intestinal side effects after metformin administration (before (T) and after (Ta) taking metformin, respectively). Six samples from three patients who had intestinal side effects after metformin administration (before (TS) and after (TSa) taking metformin, respectively). Six samples from six health subjects (N). Genomic DNA from human stool samples clinically collected was extracted by a modified CTAB method [15]. DNA concentration, purity was monitored and was diluted to proper concentration.

PCR amplification of 16S rRNA V1-V9 and high-throughput sequencing

The full V1-V9 region of the bacterial 16S rRNA gene was amplified using the universal primer set 27F and 1492R with Barcode by using third-generation sequencing [16]. The PCR products were mixed and purified. The sequencing library was generated, assessed, and sequenced on the PacBio Sequel platform using standard protocols [17].

Processing of sequencing data

The original sequences were registered in the NCBI SRA database (registration number: PRJNA725340). The clean reads were acquired by removing the barcodes and primers, low-quality reads, and chimera sequences from raw data [18, 19]. Sequences with $\geq 97\%$ similarity were assigned to the same OTUs by Uparse software (Uparse v7.0.1001) [20]. The representative sequence for each OTU was screened for further annotation. The taxonomic information for each representative sequence was annotated by the SSUrRNA Database [21] of Silva Database [22] based on the Mothur algorithm. Alpha diversity such as Chao1, Shannon index were calculated with QIIME (Version1.9.1) and displayed with R software (Version 2.15.3). The Chao1 index was selected to identify community richness, and the Shannon index was used to identify community diversity. Tukey and Wilcox's tests were used to analyze the differences between groups, $p<0.05$ was considered statistically significant. Principal Coordinate Analysis (PCoA) based on unweighted unifrac distance calculated by QIIME software (Version 1.9.1) and displayed by WGCNA package, stat packages, and ggplot2 package in R software (Version 2.15.3). The significantly bacterial taxa between groups were identified by the linear discriminate analysis (LDA) effect size (LEfSe) method with a LDA threshold value of 2.0$^{18}$ and MetaStats at 95% confidence interval, simultaneously. Potential functional contributions of the observed microbes were inferred using PICRUSt2 [23]. Significantly different Kyoto Encyclopedia of Genes and Genomes (KEGG) orthologs (KOs) were identified by t-test, $P<0.05$ was considered statistically significant. The corresponding enrichment pathways of significantly different KOs were predicted in KEGG[https://www.kegg.jp/kegg/tool/map_pathway2.html].

Bioinformatics analysis

In order to explore the effects of intolerance-related differential bacteria and primary bile acid on the body, we used the GEO database to search the research on them. As a result, we found the GSE23630 data set
related to *Lactobacillus plantarum* 299v the GSE41734 data related to *Lactobacillus brevis* 119-2, the GSE55443 data related to cholic acid. Besides, using related scores >1.0 as the cutoff we searched metformin intolerance symptoms, including bloating, diarrhea, and nausea in GeneCards (https://www.genecards.org/) to collect intolerance symptoms related genes.

Next, we used GEO2R to analyze the above two data sets related to the differential bacteria and carry out the T-test separately. Using P<0.05 as the cutoff, genes significantly different between the phorbol 12-myristate 13-acetate (PMA)/ionomycin (IO)-induced intestinal explants pro-inflammatory disease model and *Lactobacillus plantarum* 299v treated samples, and between livers samples from *Lactobacillus brevis* 119-2 and control diet-administrated rat were filtered out. If there are multiple transcripts for the same gene, we multiplied the P-value and used the square root of the product as the final P-value. Because there is only 1 sample in the case and control groups in the GSE55443 data set, we could not carry out statistical analysis. So, we selected the top 100 different genes with the maximum value and the top 100 genes with the minimum value of LogFC between CA and vehicle-treated intestinal epithelial cells of mice. Then, we intersected the differential genes obtained from the above analysis with the genes related to intolerance symptoms.

After that, protein interaction analysis on differential genes from the above analysis, and seven genes related to metformin glycemic response, and four genes related to metformin intolerance retrieved from the literature [24] was performed by String (https://string-db.org/) [25]. The interaction results were visualized by Cytoscape [26].

**RNA Isolation and cDNA Synthesis**

Total RNA was isolated from a 1ml whole blood sample with Trizol (Invitrogen, USA) reagent and purified using RNA simple Total RNA Kit (TIANGEN, China) followed the manufacturer's instructions. About 0.2μg of total RNA was used for first-strand cDNA synthesis by using Mix in FastKing RT Kit (With gDNase) (TIANGEN, China) according to the manufacturer's instructions.

**Primer Design and Evaluation**

The primer pairs of *FOXA2, GLI1, HTR7*, and *GAPDH* were designed according to their sequences by using the online program Primer-BLAST (Table S9) (https://www.ncbi.nlm.nih.gov/tools/primer-blast/ and https://pga.mgh.harvard.edu/primerbank/index.html). The primer evaluation was carried out according to the reported method of literature [27] with modification. Standard curves of each primer pair were established using 8-fold dilution series of template cDNA.

**Real-Time PCR**

The quantitative real-time PCR was performed using FastStart Essential DNA Green Master (Roche, Switzerland) with a Real-Time PCR System (BIO-RAD, USA). Each reaction contained 5 μL cDNA (8-time diluted), 10 μL FastStart Essential DNA Green Master, 1 μL Forward Primer (10 uM), 1 μL Reverse Primer
(10 uM), and 3µL RNase-Free ddH2O water. The PCR was carried out as the following steps:
predegeneration at 95°C for 10 min; 45 cycles of degeneration at 95°C for 15 s, annealing at 60°C for 15 s,
and extension at 72°C for 30 s; and melting curve analysis at 65°C-95°C. RT-qPCR of each cDNA sample
was carried out three times as technical replicates. Finally, the relative expression was determined by
using the $2^{\Delta\Delta Ct}$ method [28]. The Ct-value (cycle threshold) determined at the end of the reaction
indicates the cycle number at which fluorescence passes a fixed threshold. The amplification of β-Actin
was performed as control and reference. The differences between groups were tested by the Paired t-test.
P ≤ 0.05 and 0.05 < P ≤ 0.1 are considered significant and trend, respectively.

Abbreviations
T2D: type 2 diabetes; GI: gastrointestinal; PBA: primary bile acids; SBA: secondary bile acid; CA: cholic
acid; KOs: Kyoto Encyclopedia of Genes and Genomes (KEGG) orthologs; IBD: inflammatory bowel
disease; IGT: impaired glucose tolerance; OGTT: oral glucose tolerance test; LEfSe: linear discriminate
analysis (LDA) effect size. PcoA: Principal coordinate analysis; PMA: phorbol 12-myristate 13-acetate ;
IO: ionomycin; T: T2D individuals before metformin administration who did not have metformin-related GI
intolerance after metformin taking; TS: T2D individuals before metformin administration who had
metformin-related GI intolerance after metformin taking; Ta: T2D individuals after metformin
administration who did not have metformin-related GI intolerance after metformin taking; TSa: T2D
individuals after metformin administration who had metformin-related GI intolerance after metformin
taking.

Declarations

Ethics approval and consent to participate
The research activities were approved by the local ethics committee of First People's Hospital of Qujing
City. All clinical data collection and genetic diagnoses were performed after obtaining consent from the
patients. The objective, materials, and methods of this research and the rights and obligations of the
patients have informed patients in oral and written form. Make sure the parents understand all
information, then the informed consent was signed before study inclusion. All informed consents were
collected and delivered to the study coordinators by the doctor concerned.

Consent for publication
Not applicable.

Availability of data and materials
The original sequences were registered in the NCBI SRA database (PRJNA725340). Other data generated or analyzed during this study are included in this article and its additional files.

**Competing interests**

The authors declare no conflict of interest.

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**Authors' contributions**

Huixiang Zhang and Xue Cao designed the research, analyzed the data, wrote the main manuscript and revised the draft of the manuscript writing. Jiao Lai, Wei Zhang and Weiwen Chen collected the fecal samples and helped to write the manuscript writing. Xun Liu, Limei Wang and Lihuan Zhang collected data from databases and published studies, performed the experiments and prepared figures and tables of this paper. Qilin Gong, Rui Zhao, Dongqing Li, Hehua Huang and Ya Zhao helped to collect the fecal samples and write the manuscript. Shan Yan, Ming Yu, Xiaodan Liu and Lan-Lan Shi helped to perform the experiments and gave some advice on data analysis.

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**Figures**
Figure 1

Comparison of intestinal flora before and after taking metformin between people who had intestinal side effects and those without intestinal side effects after taking metformin, and the interaction analysis of the differential flora and side effects related genes. a-d. Relative abundance of differential intestinal flora among four groups. b-c. Principal coordinate analysis (PCoA) and linear discriminant analysis (LDA) effect size (LEfSe) analysis among groups. The red represents before taking metformin, and the green...
represents after taking metformin. e. Interaction analysis of the glycemic response and intolerance-related genes reported in the literature, differential intestinal flora, and Cholic acid. CA Cholic acid, CALP Cholic acid and Lactobacillus plantarum, SEG side effects related genes, GRG glycemic response-related genes, LB Lactobacillus brevis, LBLP Lactobacillus brevis, and Lactobacillus plantarum.

**Figure 2**

Diagram of the possible gut microbial mechanism of intestinal side effects caused by metformin. The black arrow indicates the result of the GEO analysis and RT-qPCR test, and the blue arrow indicates the results from the literature analysis.

**Supplementary Files**

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- FigureS1.pdf
