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Isomaltulose alleviates acute colitis *via* modulating gut microbiota and the Treg/Th17 balance in mice†

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Food-grade isomaltulose exhibits significant modulation of gut microbiota and its metabolites in healthy populations. This study further explored the preventive therapeutic effect and anti-colitis potential of isomaltulose on dextran sulfate sodium-induced colitis in mice. Our results suggested that isomaltulose played a significant role in preventing colon shortening, reducing intestinal epithelial destruction and inhibiting inflammatory cell infiltration. Meanwhile, the isomaltulose supplement greatly reduced the production of proinflammatory cytokines and restored the balance between T helper type 17 (Th17) cells and regulatory T (Treg) cells. Pathway enrichment analysis for differentially expressed genes (DEGs) also indicated that the anti-inflammatory effect of isomaltulose was closely related to intestinal immunity. Moreover, the disturbed gut microbiota in ulcerative colitis (UC) was partially restored after treatment with isomaltulose. These results suggest that isomaltulose is a promising therapeutic agent for the prevention and adjunctive treatment of UC by maintaining intestinal immune homeostasis and remodeling the gut microbiota.

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1. Introduction

Ulcerative colitis (UC) is a chronic recurrent inflammatory disease of the intestine involving various complications, including recurrent diarrhea, bloody stools, and tenesmus. Although its etiology is unclear, UC is closely related to the interaction between environmental factors, genetic susceptibility, and the immune system. Depending on the severity and course of UC, the therapeutics typically involve 5-aminosalicylic acid, corticosteroids and thiopurines, anti-TNF, anti-IL-12/23 p40 and anti-integrin or JAK inhibitors. These pharmacological treatments only alleviate symptomatic relief, not cure, and increase the risk of serious infection, malignant tumors and thrombosis. In recent years, gut microecological therapeutic strategies (e.g., probiotics and prebiotics) have played a greater role in the control and prevention of UC.

Gut microbiota contains the largest reservoir of the human microbiome, providing metabolic, immunologic, and protective functions for human health.^{6,7} The integrity of the intesti-

nal mucosal barrier is related to its interactions with the gut microbial ecosystem. In UC patients, the impaired intestinal epithelial barrier allows the passage of harmful gut microbiota, which activates immune cells and exacerbates the symptoms of UC. Prebiotics, as powerful modulators of gut microbiota, can directly or indirectly promote the upregulation and relocation of interepithelial tight junction proteins that form the microscopic scaffolding of the gut barrier, thereby repairing the damaged intestinal barrier in UC.⁷

Isomaltulose, composed of α -1,6-linked glucose and fructose, is naturally present in sugarcane or honey, and has been widely used as a sugar substitute in the food industry. Recent studies have reported its low glycemic index and non-carcinogenicity; thus isomaltulose was simultaneously used to improve insulin response, control hyperglycemia and reduce fat. Isomaltulose modulates the activity of specific probiotic bacteria and influences short-chain fatty acid (SCFA) production in *in vitro* co-culture systems. In addition, isomaltulose exerted beneficial effects on the regulation of gut microbiota by promoting the growth of beneficial bacteria, reducing harmful bacteria, and increasing the secretion of secondary bile acids and SCFAs in rats. In

In this study, we investigated the preventive effect of isomaltulose on dextran sulfate sodium (DSS)-induced acute colitis in mice. Our results suggested that isomaltulose significantly improved the preventative efficacy, promoted the colonic epithelial barrier integrity, modulated gut microbiota and ameliorated the regulatory T (Treg)/T helper type 17 (Th17) balance.

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2. Materials and methods

2.1. Materials and supplies

The preparation and determination protocol of isomaltulose is according to our previous method.¹³ Antibodies against occludin (A2601) and β-actin (AC026) were purchased from ABclonal (Wuhan, China). Antibodies against zonula occludens-1 (ZO-1, A11417) were purchased from Servicebio Technology Co., Ltd (Wuhan, China). Antibodies against CD3 (100203), CD4 (100407), CD25 (102010), IL-17A (506915), Foxp3 (126407) and a mixture containing phorbol 12-myristate 13-acetate (PMA), ionomycin, Brefeldin A, *etc.* (423303, BioLegend, San Diego, CA, USA) were obtained from BioLegend (San Diego, CA, USA). DSS for the colitis model was purchased from MP Biomedicals (molecular weight: 36–50 kDa, Santa Ana, CA, USA). The experimental mice and standard rodent chow food were purchased from Jinan Pengyue Laboratory Animal Breeding Company (Jinan, China).

2.2. Animal experiments

All animal procedures were performed in accordance with the Guidelines for Care and Use of Laboratory Animals of Qingdao University and experiments were approved by the Animal Ethics Committee of Medical College of Qingdao University (no. QDU-AEC-2022314). After 1 week of acclimation, C57BL/6J mice (18–20 g) were randomly grouped (n=6) as shown in Fig. 1B: (1) NC group: no treatment for all 3 weeks; (2) DSS group: DSS-induced colitis in the last week; (3) IsoMTL group: isomaltulose (400 mg per kg per day) for all 3 weeks, and DSS-induced colitis in the last week.

Acute colitis was induced by free drinking water containing 2.5% DSS. The body weight of the mice was recorded daily. Fecal samples were collected before the end of the experiment and stored at $-80~^{\circ}\text{C}$ under liquid nitrogen for gut microbiota analysis. After the mice were anesthetized with chloral hydrate, the eyeballs were removed for blood sampling. Then, the serum was obtained by centrifugation (3500 rpm, 4 $^{\circ}\text{C}$, 30 min). Under sterile conditions, the spleen tissues of mice were collected for flow cytometry analysis. The entire colon was resected, and the length of the colon was measured. The collected serum samples and colon tissues were stored at $-80~^{\circ}\text{C}$ for further analysis.

2.3. Flow cytometry analysis

Primary spleen cells were extracted, diluted and counted for flow cytometry analysis (BD FACSVerseTM, NJ, USA).¹⁴ To determine the number of Th17 cells, collected cells were pre-treated with the PMA mixture. For Th17 and Treg cell counts, CD3, CD4 and CD25 were used as surface markers, and IL-17A and Foxp3 were used as intracellular/intranuclear cytokines, respectively.

2.4. Histopathological analysis

The collected tissues were dissected, then washed with saline three times and fixed with 10% buffered formalin (pH 7.2) solution overnight. Subsequently, the tissues were rung cut and placed on a glass slide and stained with hematoxylin and eosin (H&E) and alcian blue by Servicebio Technology Co., Ltd (Wuhan, China). The images were captured using a microscope with 200× magnification (E100, Nikon, Tokyo, Japan) and an imaging system (DS-U3, Nikon, Tokyo, Japan). Mucusproducing goblet cells were observed and counted with a light microscope (OLYMPUS, Tokyo, Japan).

2.5. RNA extraction and quantitative real time PCR (RT-qPCR) analysis

The extraction of total RNA and reverse transcription were performed according to the instructions of the SparkJade kit (Jinan, China). The RT-qPCR primers are shown in Table S1.† The gene expression levels were normalized using GAPDH, and the relative quantification of gene expression was calculated by the $2^{-\Delta\Delta Ct}$ method.

2.6. Western blot and ELISA assay

Total protein of colon tissue samples was extracted using a cold RIPA lysis buffer (Solarbio, Beijing, China). Western blot analysis was performed as previously described. The ELISA kits were purchased from Nanjing Jiancheng Bioengineering Institute (Nanjing, China). The levels of proinflammatory cytokines in the serum were measured according to the instructions of the ELISA kit.

2.7. Transcriptome analysis

The transcriptome analysis was performed by Berry Genomics Co., Ltd (Beijing, China). In brief, the total RNA was extracted from colon tissues and the RNA library was constructed. The quantitative library was analyzed by single-end sequencing on the Illumina Genome analyzer. Data analysis was processed using R software. The raw data of counts and FPKM was available (https://bioinfogo.org/ResKsumo/data). The Gene Ontology (GO) and Kyoto Encyclopedia of Genes and Genomes (KEGG) analyses were performed using the "clusterProfiler", "GOplot" package from R software for the differential expression genes (DEGs), which selected the standard fold change less than or equal to 2 and the *P* value less than 0.05.

2.8. Gut microbiota analysis

Fecal DNA extraction and dilution were performed as previously described.¹⁵ The PCR was used to amplify the V3–V4 region of the bacterial 16S rRNA gene. The sequencing library was generated and sequenced using the NovaSeq 6000 platform (Shenzhen, Guangzhou, China). The following analysis of the gut microbiota structure and distribution was performed according to our previous studiy.¹⁶

2.9. Statistical analysis

Data were presented as mean \pm SD. Statistical significance between two groups was calculated using Student's t-test and that between multiple groups was determined using one-way analysis of variance (ANOVA). All statistical analyses were calculated using GraphPad Prism 9.3 (La Jolla, CA, USA).

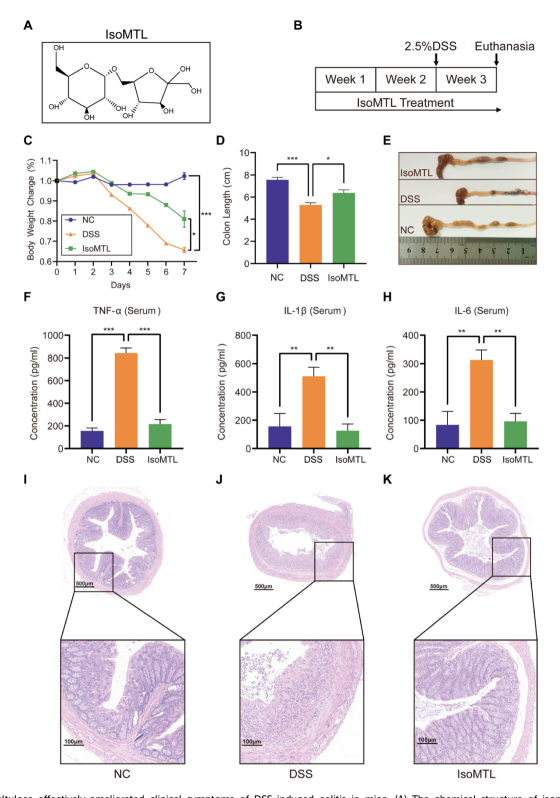


Fig. 1 Isomaltulose effectively ameliorated clinical symptoms of DSS-induced colitis in mice. (A) The chemical structure of isomaltulose. (B) Experimental design flow diagram. IsoMTL treatment: isomaltulose (400 mg per kg per day). (C) The body weight changes (n = 6). (D and E) Length of colons. The levels of TNF- α (F), IL-1 β (G) and IL-6 (H) in serum (n = 6) were determined using ELISA. (I–K) H&E-stained colon tissue. Compared to the indicated group, *P < 0.05, **P < 0.01 and ***P < 0.001.

3. Results

3.1. Isomaltulose effectively improves clinical symptoms of acute colitis in mice

Isomaltulose is an isomer of sucrose (Fig. 1A), which was prepared by isomerisation. Briefly, the sucrose isomerase gene from Pantoea dispersa UQ68J was overexpressed in non-pathogenic yeast Yarrowia lipolytica and the sucrose isomerase was used to isomerize sucrose to produce high yields of isomaltulose. Further removal of monosaccharide by-products by converting into intracellular lipids resulted in isomaltulose with a purity of up to 97.8%. 13 A DSS-induced acute colitis mouse model was used for investigating the preventive and protective effects of isomaltulose (Fig. 1B) according to our previous method.¹⁵ As shown in Fig. 1C, the body weights of mice in the DSS group decreased significantly within 7 days of DSS treatment (P < 0.001) and administration of isomaltulose (400 mg per kg per day) can significantly inhibit the weight loss caused by DSS (P < 0.05). Colon length is often evaluated as a key macroscopic indicator of colitis and is regarded as inversely associated with the severity of colitis.¹⁷ Consistent with this the colon length in the DSS group was obviously shorter (P < 0.001), which could be significantly reversed by isomaltulose (P < 0.05, Fig. 1D and E). These results indicated that isomaltulose supplementation significantly reduces the DSS-induced colitis symptoms. The pro-inflammatory cytokines (e.g. TNF-α, IL-1β and IL-6) commonly decide the onset and progression of inflammation.¹⁸ To elucidate the effect of isomaltulose on the inflammatory response, the levels of proinflammatory cytokines in serum were measured (Fig. 1F-H). Compared with NC group, TNF-a, IL-1 and IL-6 in the DSS group were significantly increased. Interestingly, isomaltulose supplementation significantly reduced the levels of these proinflammatory cytokines. These results suggested that the administration of isomaltulose significantly inhibited the expression of pro-inflammatory cytokines and reduced the inflammatory response in a DSS-induced colitis mouse model.

To further explore the protective effect of isomaltulose on the colon, histological staining and examination of colon sections from mice were performed (Fig. 1I–K). Compared with the NC group, the histopathological examination of the DSS group showed infiltration of inflammatory cells, the absence of glands, mucosal epithelial necrosis (Fig. 1I and J). In contrast, isomaltulose supplementation significantly alleviated the severe histological damage induced by DSS, as manifested by the increased glandular numbers, mucosal epithelial integrity, and reduced inflammatory cell infiltration (Fig. 1K). Based on the above results, isomaltulose has a good protective effect on ameliorating the pathological damage of the colon caused by DSS.

3.2. Effect of isomaltulose on the integrity of the intestinal barrier

The major tight junction proteins ZO-1, occludin, and claudin-1 of the intestinal epithelium involved in mucosal healing form an important barrier in the gut, preventing the spread of potentially harmful pathogens and toxins¹⁹ (Fig. 2A). Therefore, to determine the regulatory role of isomaltulose in intestinal tight junction proteins, we measured the expression of major tight junction proteins using western blot (Fig. 2B) and RT-qPCR (Fig. 2C–E). As shown in Fig. 2B, isomaltulose significantly increased the protein expression of ZO-1 (P < 0.01) and occludin (P < 0.05). Meanwhile, isomaltulose restored the gene transcript levels of ZO-1 (P < 0.01), occludin (P < 0.001) and claudin-1 (P < 0.01) that were decreased by DSS (Fig. 2C–E).

The mucus layer protects the intestine from the external environment, including pathogens.²⁰ Alcian blue staining was further used to determine the mucosal barrier function (Fig. 2F–H). The number of mucus-producing goblet cells in the colons of mice administered with isomaltulose was significantly increased compared to mice in the DSS group. All these results suggested that isomaltulose treatment had a protective effect on the integrity of the intestinal barrier in mice with DSS-induced colitis.

3.3. Effect of isomaltulose on the intestinal immune response

To investigate the mechanism by which isomaltulose alleviates colitis, we performed transcriptome analysis. As shown in Fig. 3A, the X-axis log 2 (fold change) represents the differential expression fold and up-regulation or not of DEGs is judged according to whether it is a positive value or not. Genes with larger differences were distributed closer to the two ends of the X-axis, which were the red and green dot parts. The Y-axis is represented by the -log 10 p-value, which is inversely proportional to the p-value and proportional to the significance of the degree of difference. A total of 984 DEGs were obtained, including 739 upregulated and 245 downregulated genes (Fig. 3A). Furthermore, isomaltulose treatment significantly reversed the gene expression pattern of mice in DSS-induced colitis mice (Fig. 3B). GO and KEGG enrichment analyses of DEGs were performed to better understand the underlying molecular mechanism and function of isomaltulose. Most of the GO terms of the biological process (BP) were focused on the mononuclear cell proliferation, immune system processes, leukocyte cell-cell adhesion, and T cell activation (Fig. 3C). The KEGG pathway analysis for the top 20 immune related pathways showed that DEGs were involved in pathways such as the B cell receptor signaling pathway, cytokine-cytokine receptor interaction, IL-17 signaling pathway, TNF signaling pathway, T cell receptor signaling pathway and PI3K-Akt signaling pathway (Fig. 3D). In particular, enrichment in the IL-17 signaling pathway and T cell receptor signaling pathway suggests that the role of isomaltulose has an important relationship with Th17-mediated immunity.

3.4. Isomaltulose improves the balance of Treg/Th17 cells

IL-10 is the main product of Treg cells and plays a key role in Treg cell-mediated remission of colitis.²¹ IL-17 is the main effector of Th17 cells,²² whose family includes six-member ligands (IL-17A–IL-17F) and 5 receptors (IL-17RA–IL-17RD and

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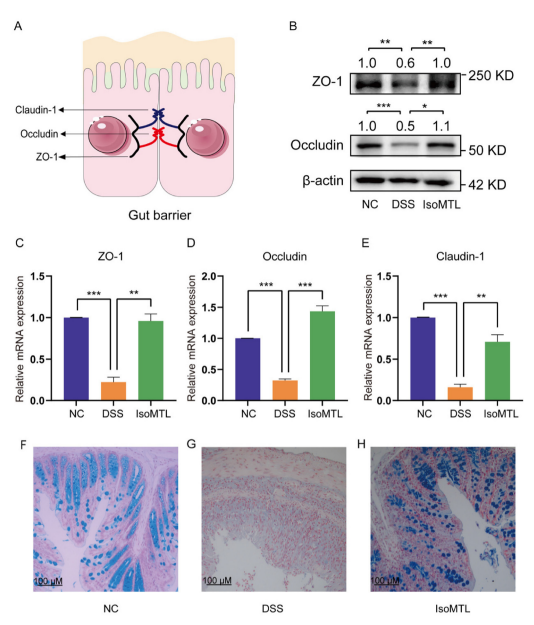


Fig. 2 Effects of isomaltulose on intestinal tight junction protein expression. (A) Cell tight junction pattern. (B) Western blot analysis of occludin and ZO-1 (n = 3). (C-E) The relative mRNA expression of ZO-1, occludin and claudin-1. (F-H) Representative images of alcian blue staining of colon sections. Compared with indicated groups, *P < 0.05, **P < 0.01 and ***P < 0.001.

SEF), ²³ and the pathogenic role of IL-17A has been well documented in UC. ²⁴ Transcriptomic analysis of the Th17 and Treg related signaling pathways showed that DSS induction could upregulate the gene expression of the Th17 related signaling pathway and reduce the gene expression of the Treg related signaling pathway. Administration of isomaltulose can reverse the expression pattern of genes in Th17 and Treg signaling pathways (Fig. 4A). Specifically, the gene expression of IL-17A, IL-17F and IL-17RB was significantly decreased and the main effector of Treg cells Foxp3 ²² was highly expressed in the IsoMTL group. In addition, IL-2 inhibited Th17 differentiation ²⁵ and promoted the differentiation and function of Treg cells, ²⁶ which was significantly expressed in the IsoMTL

group. Therefore, this result indicated that isomaltulose can regulate the balance of Treg/Th17 cells at the gene level. To further explore the function of isomaltulose on the balance of Treg/Th17 cells, we used RT-qPCR (Fig. 4B and C) and flow cytometry (Fig. 4D and E) to detect changes in cytokine production and Treg/Th17 cell differentiation. Compared to the NC group, the DSS treatment group significantly increased the expression of IL-17 (P < 0.001) and decreased the expression of IL-10 (P < 0.05). Compared with the DSS group, isomaltulose supplementation significantly increased the expression of IL-10 (P < 0.01), while greatly decreasing the expression of IL-17 (P < 0.01) in colitis mice (Fig. 4B and C). Consistent with this isomaltulose significantly decreased the number of Th17

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A В Filtered Up 12 Down 10 -Log10 (P Value) 8 6 2 Log2 (Fold Change IsoMTL vs DSS) DSS D C Top 30 of Enrichment Top 20 of Enrichment structural constituent of ribosome Cytokine-cytokine receptor interaction cytosolic ribosome Viral protein interaction with cytokine ribosomal subunit Chemokine signaling pathway respirasome mitochondrial protein-containing complex Cell adhesion molecules(CAMs) cytosolic small ribosomal subunit B cell receptor signaling pathway negative regulation of immune system process mitochondrial respirasome cytosolic large ribosomal subunit mitochondrial membrane protein complex Leukocyte transendothelial migration IL-17 signaling pathway positive regulation of response to exte Jak-Stat singaling pathway mononuclear cell prolif VEGF singaling pathway small ribosomal subunit respiratory chain complex Rap1 singaling pathway leukocyte proliferation leukocyte cell-cell adhesion HIF-1 singaling pathway regulation of T cell activation TNF singaling pathway T cell singaling pathway lymphocyte proliferation large ribosomal subunit PI3K-Akt singaling pathway regulation of mononuclear cell proliferation regulation of leukocyte proliferation ErbB singaling pathway immune receptor activity oxidoreduction-driven active transmembrane transporter activity negative regulation of leukocyte activation regulation of cell-cell adhesion MAPK singaling pathway C-type lectin receptor singaling Calcium singaling pathway leukocyte chemotaxis regulation of lymphocyte proliferation Primary immunodeficiency regulation of leukocyte cell-cell adhesion Inflammatory mediator regulation of TRP channels 12 2.2 2.8 3.4 4.0 Enrichment_score **Enrichment score** P-value GeneNumber 3

Fig. 3 Enrichment analysis and deferentially expressed genes. (A) Volcano map of DEGs between isomaltulose and DSS. Red represents up-regulated genes and green represents down-regulated genes. (B) The expression profiles of DEGs. The transition from high to low expression is represented by the red to blue colors. (C) Top 30 enrichment was displayed using GO analysis. (D) KEGG pathway enrichment analysis of top 20 immune related pathways. GO and KEGG enrichment analysis were selected according to the rank of q-value.

0.3

0.1

cells (P < 0.01, Fig. 4D) and increased the number of Treg cells in the spleen of the DSS-induced colitis model (P < 0.05, Fig. 4E). These results indicated that isomaltulose could restore intestinal immune homeostasis in the mouse model of UC by rebalancing the Treg/Th17 cell populations and the production of pro-inflammatory and anti-inflammatory cytokines.

3.5. Isomaltulose alters the relative abundance of gut microbiota

The regulatory effect of isomaltulose on gut microbiota was further studied by 16S rRNA sequencing. The common and unique OTUs were displayed to prove that DSS induction and

2e-14

1e-14

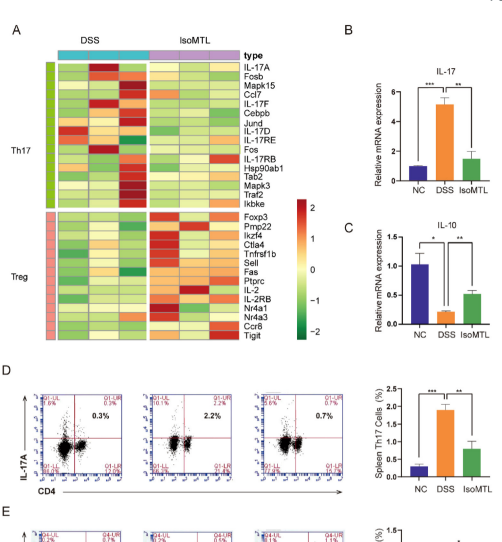


Fig. 4 Isomaltulose improves the DSS-disturbed balance between Th17 and Treg cells. (A) Expression pattern of DEGs in Th17 and Treg signaling pathways. The transition from high to low expression is represented by the red to green colors. The mRNA expression of IL-17 (B) and IL-10 (C) in colon tissues. Flow cytometry analysis of (D) CD3+CD4+IL-17A+ Th17 cells and (E) CD3+CD4+CD25+FoxP3+ Treg cells. *P < 0.05, **P < 0.01, ***P < 0.001.

isomaltulose treatment changed the composition of gut microbiota (Fig. 5A). The microbiota composition in each group was clearly clustered using weighted UniFrac principal coordinates analysis (PCoA). In PCoA analysis, the percentages presented along the axes represent the overall fraction of dissimilarity captured by PC1 (50.1%) and PC2 (29.42%) (Fig. 5B). Compared to the NC group, the DSS administration significantly shifted the structure of gut microbiota and isomaltulose reversed this shift to some extent. Simultaneously, the relative abundance of gut bacteria was compared at the phylum level (Fig. 5C). We observed that isomaltulose significantly reversed

CD25

0.7%

the DSS-induced changes in *Firmicutes*, *Bacteroidota*, and *Proteobacteria*. As shown in Fig. 5D–F, mice fed with DSS presented a drastic reduction in the F/B ratio, and isomaltulose treatment reduced the relative abundance of *Firmicutes* and increased the relative abundance of *Bacteroidota* caused by DSS. Compared to the DSS group, the relative abundance of *Proteobacteria* and *Actinobacteriota* was partly reversed after the administration of isomaltulose (Fig. S1A–C†). It is worth noting that the relative abundance of *Verrucomicrobiota* was significantly increased, which could increase the content of SCFAs.

NC

Spleen Treg Cells

DSS IsoMTL

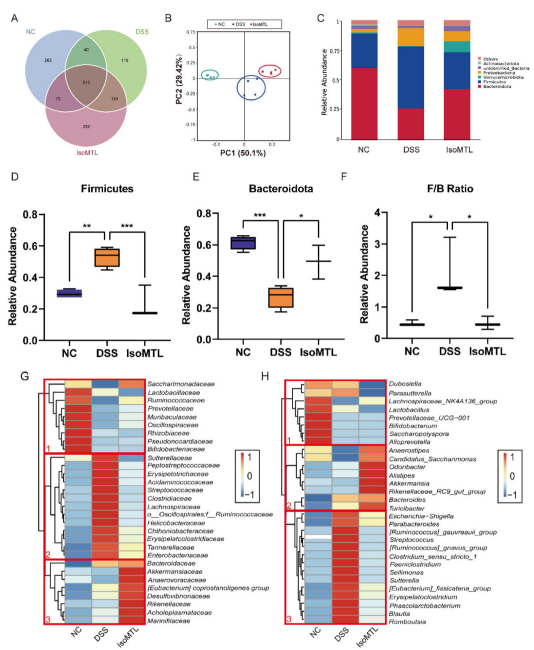


Fig. 5 Isomaltulose modulated the overall structure of the gut microbiota. (A) Common and unique OTUs. (B) Weighted UniFrac PCoA analysis. (C) The phylum-level organization of the gut microbiota. Relative abundance of *Firmicutes* (D), *Bacteroidetes* (E), F/B ratio (F) in the three groups. Microbial community species abundance heatmaps at the family level (G) and genus level (H). The species abundance heatmap shows the abundance distribution of the 30 families or genera (Y-axis) that are predominant in all samples (X-axis). The abundance of bacteria is indicated by color intensity.

In order to further determine the composition differences of gut microbiota in mice, the changes of gut microbiota at the level of family and genus were observed (Fig. 5G and H). The DSS group showed clear differences from the NC group at both the family and genus levels, indicating that the DSS-induced colitis mice caused obvious gut dysbiosis. At the family level, the gut microbiota of cluster 2 increased after DSS treatment, while the gut microbiota of cluster 1 decreased (Fig. 5G). For cluster 1, the abundance of *Bifidobacteriaceae*,

Pseudonocardiaceae, and Lactobacillaceae was higher in the NC group than in the DSS and IsoMTL groups. After the treatment with isomaltulose, the changes of gut microbiota in cluster 2 were obviously reversed, and the bacteria were partially restored to the level of the NC group. Meanwhile, the gut microbiota of cluster 3 was obviously elevated under the effect of isomaltulose, which mainly consisted of beneficial bacteria (Akkermansiaceae, Marinifilaceae, Anaerovoracaceae, etc.). At the genus level, the gut microbiota in cluster 1 was decreased and

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that in cluster 3 was increased after DSS treatment (Fig. 5H). After isomaltulose treatment, the gut microbiota of cluster 3 was restored to the control level, and the gut microbiota of cluster 2 was significantly increased (e.g., Akkermansia and Alistipes). These results further suggested that isomaltulose might reduce the effect of DSS on gut microbiota by increasing partially the content of beneficial bacteria, as well as restoring the dysbiosis of gut microbiota.

The LEfSe method was used to analyze the difference between the samples, which could further show the microbial diversity (Fig. 6A and B). The larger the LDA score, the greater the influence of species abundance on the difference effect. Setting LDA score ≥ 2 and P < 0.05 as a meaningful cut-off, LEfSe identified 32 bacterial taxa which were significantly different between the three groups and could represent typical characteristics of different groups. As Fig. 6A shows, the DSS group had 15 dominant microorganisms, while the IsoMTL group had the other 9 dominant microorganisms, including families Bacteroidaceae, genus Bacteroides and phylum Gemmatimonadetes, etc., which was also verified by the differential-abundance microbial cladogram (Fig. 6B). Thus, LEfSe analysis illustrated that multiple taxonomic differences were found among three groups of DSS, NC and IsoMTL.

3.6 Association between gut microbiota and UC-related symptoms and Treg/Th17-related immunity

Correlation analysis can be used to verify the correlation between different data, so we used Spearman correlation analysis to examine the relationship between inflammatory factors and gut microbiota. Based on our results, inflammatory factors (TNF-α, IL-6, IL-17 and IL-1β) and tight junction pro-

teins (ZO-1 and occludin) were highly correlated with changes in gut microbiota. As shown in Fig. 7, genus of Peptostreptococcaceae, Romboutsia, Blautia, Streptococcus, Lachnospiraceae and Ruminococcaceae showed a significantly positive correlation with inflammation factors and displayed a negative correlation with tight junction proteins. Meanwhile, the presence of Enterobacteriaceae and Parabacteroides significantly reduced the body weight and colon length. Notably, Akkermansiaceae and Alistipes were significantly positively correlated with Treg cells, while Parasutterella was significantly negatively correlated. Besides, there was a significantly positive correlation between the abundance of Bacteroidaceae and Bacteroides with the level of iNOS. These results clearly reveal the relationship between changes in different bacteria and host biological effects, indicating that changes in the gut microbiota are closely associated with inflammatory responses, especially tight junction proteins.

Discussion 4.

Isomaltulose is an isomer of sucrose²⁷ derived from natural sources such as honey (<1%) and sugarcane, 28 whose sweetness is approximately half that of sucrose (0.4-0.45).²⁹ We previously developed engineered yeast to obtain high-purity isomaltulose. 13 In healthy and obese patients, isomaltulose supplementation had been reported to reduce insulin resistance and elevated blood pressure, while benefiting endothelial function and cardiovascular health. 30,31 Herein, we further indicated that isomaltulose supplementation was significantly effective in improving the prevention of acute colitis in mice, promoting colonic epithelial barrier integrity, reducing inflam-

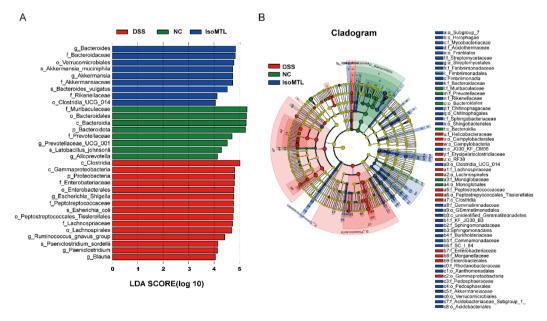


Fig. 6 Alteration of the gut microbiota by isomaltulose. (A) Linear discriminant analysis (LDA) effect size (LEfSe) and (B) cladogram of species annotated by mOTU were used to identify differentially abundant taxa within the DSS, NC, and IsoMTL groups. P < 0.05 and LDA score (log 10) > 2 was considered significant; (B) cladogram displayed the taxonomic tree of differentially abundant taxa by LEfSe analysis.

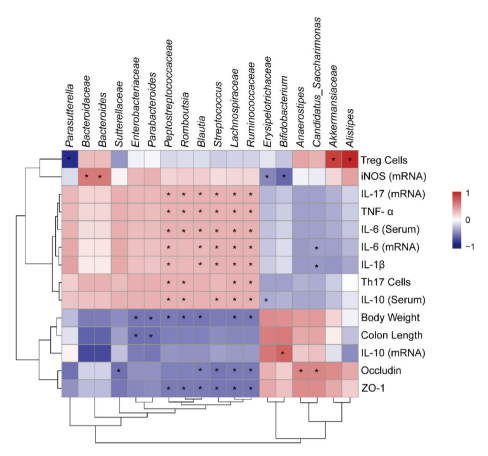


Fig. 7 Correlation analysis of UC-related symptoms and Treg/Th17-related immunity with the gut microbiota. Correlation heatmap calculated by Spearman's correlation coefficient. Colors range from blue (negative correlation) to red (positive correlation), *P < 0.05, significant correlation.

matory cytokines, modulating gut microbiota and ameliorating Treg/Th17 balance (Fig. 8). These results suggest that isomaltulose is a promising therapeutic agent for the prevention and adjuvant treatment of UC by maintaining intestinal homeostasis.

The pathogenesis of UC is closely related to a compromised intestinal barrier, where hyperpermeability of the gut increases pathogen invasion and leads to translocation of the gut microbiota.32 Tight junction proteins (e.g., ZO-1, claudin-1 and occludin) play an important role in maintaining cellular integrity and regulating intestinal barrier permeability by regulating the paracellular permeability of water, ions and macromolecules in adjacent cells.33 Reduced expression of intestinal tight junction proteins can cause local or systemic inflammatory and immune responses in the gut, as well as endotoxin invasion, intestinal bacterial translocation and disruption of the intestinal mucosal barrier, which can further exacerbate local or systemic inflammatory and immune responses in the gut.34-37 Previous studies have identified the role of isomaltulose in reduction of intestinal pathogens. 12 However, only a few studies have focused on the effects of isomaltulose on tight junctions and related intestinal barrier. In this study, morphology (Fig. 1I-J), transcript and protein analysis (Fig. 2) all demonstrated the excellent therapeutic role of isomaltulose

in improving the intestinal barrier and tight junctions disrupted by DSS, further explaining the action model of its role in UC adjuvant therapy.

Recently, isomaltulose has been shown to alter the structure and composition of the gut microbiota, thereby increasing the abundance of beneficial microorganisms, suggesting its potential prebiotic activity in the normal polulation. 12 Herein, we demonstrated that isomaltulose treatment greatly modulated the overall structural and genus/species-specific changes in the dysfunctional gut microbiota of DSS-induced colitis mice (Fig. 5 and 6). An increase in Lachnospiraceae can cause immune overresponse triggering intestinal inflammation.³⁸ Meanwhile, a study had shown that Lachnospiraceae and Streptococcaceae were increased in the gut in mice with colitis, 39 which was consistent with our results (Fig. 5G). After treatment with isomaltulose, the number of Lachnospiraceae and Streptococcaceae were significantly reduced. He et al. proposed that prebiotic unsaturated alginate oligosaccharides could improve colitis via modulating gut microbiota, such reducing the abundance of Acidaminococcaceae and Lachnospiraceae, while increasing the abundance Anaerobiospirillum, which showed a similar pattern to isomaltulose.40 Furthermore, isomaltulose treatment significantly increased the number of some anti-inflammatory bacteria

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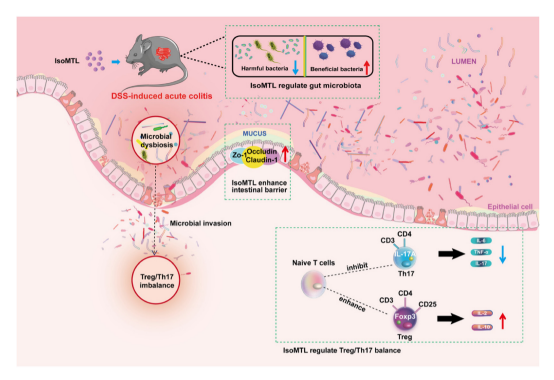


Fig. 8 Possible mechanism of isomaltulose prophylactic treatment of DSS-induced colitis.

(*e.g.*, *Bacteroides*, *etc.*) and the amount of *Akkermansia*, a typically beneficial bacteria known to be positively associated with gut integrity.⁴¹ These findings could contribute to the development of isomaltulose as a specialty health food for UC patients.

Treg/Th17 cell imbalance is a key factor in the development of UC, ⁴² hence targeting Treg/Th17 cell regulation has emerged as a potential strategy for the prevention and treatment of UC. ^{43–45} Bursts of inflammation are the main factor in the pathogenesis of UC, and the production of cytokine storms contributes directly to this process. ^{46,47} In this study, the changes of Treg and Th17 indicated that isomaltulose treatment improved the balance of Treg/Th17 cells, which has a similar immune mechanism to an active ingredient of Chinese herbal medicine used to treat UC called baicalin. ⁴⁸ Spearman analysis further showed that changes in the gut microbiota were strongly correlated with associated changes in the gut barrier, inflammatory factors and Treg/Th17 cells (Fig. 7).

5. Conclusion

In this study, a food-grade supplement of isomaltulose significantly ameliorates DSS-induced UC symptoms and gut microbiota dysbiosis in mice. Meanwhile, the changes of specific families/genus in gut microbiota were significantly associated with UC-related symptoms, tight junction proteins, inflammatory factors, and the Treg/Th17 cell balance. Our results suggest that isomaltulose is a promising therapeutic agent for UC prevention and adjuvant treatment *via* maintaining intestinal immune homeostasis and remodeling intestinal microbiota.

Abbreviations

DSS	Dextran	sodium	sulfate

ELISA Enzyme linked immunosorbent assay

H&E Hematoxylin and eosin

IL Interleukins

SCFAs Short chain fatty acids

Th17 T helper 17 cell

TNF Tumor necrosis factor

Treg T regulatory cell

ZO-1 Zonula occludens-1

DEGs Differentially expressed genes

PMA Phorbol 12-myristate 13-acetate

GO Gene ontology

KEGG Kyoto encyclopedia of genes and genomes

Author contributions

S. L. and N. H. designed the study; Z. Z., S. Y., L. C., H. P. and K. S. performed the experiments; Z. Z. and N. H. analyzed the data and made figures; Z. Z., L. C. and S. Y. wrote the manuscript. S. L., N. H., Z. Z., L. C. and S. Y. reviewed/edited the manuscript. All authors approved the final manuscript.

Conflicts of interest

The authors state no conflicts of interest.

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