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2023 Doctoral Thesis

The role of Paneth cell α -defensin as a regulator of the intestinal microbiota in nonalcoholic steatohepatitis

(非アルコール性脂肪肝炎における腸内細菌叢の制御 因子である Paneth 細胞α-defensin の役割)

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Chapter 1. General Introduction

Nonalcoholic steatohepatitis (NASH) is a chronic liver disease characterized by steatosis, liver injury, inflammation, and fibrosis¹. NASH is the crucial risk factor for cirrhosis and hepatocellular carcinoma and regarded as one of the urgent unmet medical needs with the rapidly increasing number of patients all over the world². The liver fibrosis, which is a characteristic pathology of NASH, is associated with an increased risk for mortality, so that the prevention of fibrosis is important³. Obesity and type 2 diabetes are known to relate with NASH, and overnutrition is the principal cause of these disease⁴. Because the fibrosis progression rate among nonalcoholic fatty liver disease (NAFLD) patients is about 40% and NAFLD has a very variable natural history, a multiple hit hypothesis has been considered in which NASH development proceeds through the complex crosstalk of varied factors not limited to the liver^{5,6,7}. The progression of NASH pathology is associated with overnutrition⁸, yet the pathophysiological mechanisms of NASH are not well understood⁹.

The human intestine harbors intestinal microbiota with 40 trillion bacteria, which affects health and disease via controlling host metabolism, immune function, and more^{10,11}. On the other hand, dysbiosis, disruption of intestinal microbiota homeostasis, is associated with many diseases including metabolic disorders such as fatty liver and obesity^{12,13}. The liver plays a central role in metabolism of the body and directly communicate with intestine to form the "gut-liver axis", and the intestinal microbiota has a key role in the function of liver¹⁴. The metabolites, which are produced by the intestinal bacteria, such as secondary bile acids and short-chain fatty acids (SCFA) regulate liver fatty acid synthesis and bile acid synthesis^{15,16}. Recently, the relationship between dysbiosis and NAFLD/NASH has been revealed. The composition of the intestinal microbiota in NASH patients is different from healthy individual¹⁷. Among NAFLD patients, differences in the intestinal microbiota have been reported between patients with NAFL and NASH as well as between severe and mild/moderate fibrosis^{18,19}.

patterns induce chronic inflammation in the liver^{20,21}. These reports suggest that dysbiosis contributes to NASH progression.

A monolayer of the intestinal epithelium located at the interface between intestinal lumen and the lamina propria facilitates the absorption of nutrients and prevents the invasion of pathogen and more²². Paneth cells at the bottom of small intestinal crypts secrete granules rich in α -defensins, human defensin (HD) 5 in humans and cryptdins (Crps) in mice, into the intestinal lumen and contribute to innate enteric immunity by killing pathogenic bacteria^{23,24,25,26,27}. Furthermore, α -defensin was shown to have selective bactericidal activities, killing pathogenic bacteria while eliciting less bactericidal activities against commensal bacteria²⁸. Matrix metalloproteinase 7 (MMP7)-deficient mice absent of activated Crps increase the percentage of Firmicutes, in contrast, transgenic expression of HD5 decrease the percentage of Firmicutes^{29,30}. In addition, MMP7 deficient mice impair symbiotic bacterial colonization after antibiotic treatment, indicating that α -defensin promotes commensal bacterial treas³¹. These reports suggest Paneth cell α -defensin regulate the composition of the intestinal microbiota by selective bactericidal activity.

Paneth cell dysfunction has been reported to be associated with certain diseases. Deficiency in the autophagy gene *Atg16L1*, which has been identified as susceptible gene for Crohn's disease, leads to defects in granule secretion, as well as abnormal granule morphology and reduction of granule number³². In mouse model of allogeneic hematopoietic stem cell transplantation, graft-versus-host disease (GVHD) causes reduction of Paneth cells and α -defensins, that enhance bacterial translocation and further exacerbating the disease^{33,34}. In contrast, restoration of luminal α -defensin prevented GVHD-mediated dysbiosis³⁵. Furthermore, Paneth cell of severely obese patient showed abnormal granule morphology and decreased protein level of α -defensin³⁶, suggesting that dietary habits induce Paneth cell dysfunction. Taken together, these reports showed the possibility that impaired secretion of α -defensin from Paneth cell causes disease via dysbiosis³⁷.

In this study, we hypothesize that dietary factors composed with high fat triggers abnormalities in Paneth cell α -defensin to induce dysbiosis, resulting in NASH fibrosis. Here we show in diet-induced NASH model mice, the amount of α -defensin secreted from Paneth cells decreased significantly before the onset of NASH, resulting in dysbiosis with decreased microbiota diversity followed by fibrosis leading to NASH. Furthermore, restoring luminal amount of α -defensin by intravenous administration of R-Spondin1 (R-Spo1) that regenerates Paneth cells or oral administration of α -defensin both suppress dysbiosis and ameliorates liver fibrosis. Our results show for the first time that dysbiosis caused by decrease of Paneth cell α -defensin secretion in "gut-liver axis" contributes to the onset and the progression of NASH, further providing novel insights into Paneth cell α -defensin as a potential therapeutic target to contribute to prevention and treatment of NASH.

Chapter 2. Materials and Methods

2.1 Mice

Three-week-old male C57BL/6J mice were purchased from CLEA Japan Inc. and acclimated for 3 weeks (wks) prior to be using in experiments. After acclimation, six-week-old mice were divided into two groups: SD group fed with standard diet (SD; A06071314, Research Diets Inc.) and CDAHFD group fed with choline-deficient, L-amino acid-defined, high-fat diet with 0.1% methionine (CDAHFD; A06071302, Research Diets Inc.). The mice were maintained on 12-h light/dark cycle and had free access to both food and tap water. All animal experiments in this study were conducted in accordance with the approval of the Institutional Animal Care and Use Committee of the National University Corporation at Hokkaido University in accordance with Hokkaido University Regulations of Animal Experimentation.

2.2 Histological analysis

The left lobe of liver was quickly excised and fixed in 10% neutral buffered formalin. 4 μ m sections of paraffin-embedded tissue were stained using hematoxylin and eosin (H&E) or Sirius red. The average number of inflammatory foci and hepatocellular ballooning were determined from 5 randomly selected fields (228×303 μ m²) per slide by using image analysis software, NIS-Elements D ver. 4.13 (Nikon). Hepatocellular ballooning was scored as follow: 0 (ballooned cells are absent), 1 (ballooned cells are present). Using the BZ-II analyzer (KEYENCE), at least 20 fields (272×361 μ m²) per slide were quantified for lipid accumulation area and Sirius red-positive area, respectively, and the average percentage of hepatic steatosis area and fibrosis area were evaluated. TUNEL assay was performed using the apoptosis in situ detection kit (Wako) following manufacturer's instruction. TUNEL-positive cells were quantified by counting positive nuclei in 5 randomly selected fields (118×156 μ m²) per slide by using NIS-Elements D.

2.3 Reverse transcription and quantitative PCR

DNA-free RNA was obtained from ileal or liver tissue by Dnase treatment using the RNeasy Mini Kit (QIAGEN). 0.5 µg total RNA was reverse transcribed with SuperScript VILO MasterMix (Life Technologies) by thermal cycled at 25 °C for 10 min, 42 °C for 60 min, and 85 °C for 5 min using Mastercycler EP (Eppendorf). Quantitative PCR was performed using Roche LightCycler 96 (Roche) with fluorescence-labeled locked nucleic acid (LNA) hydrolysis probes (Roche) from the Universal Probe Library (UPL) following the manufacturer's protocol. Gene expression was normalized to hypoxanthine guanine phosphoribosyl transferase-1 (Hprt1). The primer sequences are listed in Table 1.

Table 1 PCR primers

Gene	Primer sequence (5'-3')	Universal Probe Library	
Pdia3	F: TGGCCACTGTAAGAATCTGGA	#33	
	R: TGACAATATTTGGATCTTTGCTG		
Perk	F: CCTTGGTTTCATCTAGCCTCA	#106	
	R: ATCCAGGGAGGGGGATGAT		
Chop	F: GCGACAGAGCCAGAATAACA	#91	
	R: GATGCACTTCCTTCTGGAACA		
Nox2	F: GATGCACTTCCTTCTGGAACA	#20	
	R: GTGCACAGCAAAGTGATTGG		
Prdx6	F: TTTCAATAGACAGTGTTGAGGATCA	#1	
	R: CCGTGGGTGTTTCACCAT		
Atg3	F: GTACCTGACCCCGGTCCT	#78	
	R: TTGGACAGTGGTGGACTAAGTG		
Atg12	F: CATTGACTTCATCAAAAAGTTCCTT	- #49	
	R: GGCAAAGGACTGATTCACATAA		
Lath	F: CCCCACCAAGATCCCAGT	- #7	
1,50	R: CGCTCATGTTCACGTGGT		
F4/80	F: GGAGGACTTCTCCAAGCCTATT	- #42	
	R: AGGCCTCTCAGACTTCTGCTT		
Cd11b	F: CAATAGCCAGCCTCAGTGC	- #76	
	R: GAGCCCAGGGGAGAAGTG		
Cd11c	F: AGCCTCAAGACAGGACATCG	- #106	
	R: TGAATCCTGGAGGGGGATCT		
Tgfb1	F: CTGGGCACCATCCATGAC	- #68	
	R: CAGTTCTTCTCTGTGGAGCTGA		
Trailr2	F: CCCTGAGATCTGCCAGTCAT	#103	
	R: TTTCTCTGGGGGGTACAGGAA		
Bax	F: GAACCATCATGGGCTGGA	- #69	
	R: GGTCCCGAAGTAGGAGAGGA		
Crp1	F: CCAAAACACAGATGAAGAGACTAAAA	#68	
	R: GCATACCAGATCTCTCAACGATT		
HPRT-1	F: TCCTCCTCAGACCGCTTTT	#95	
	R: CCTGGTTCATCATCATCGGCTAATC		

2.4 Fecal DNA extraction

Fresh fecal samples were collected immediately after excretion, snap-frozen on dry ice, and stored at -80°C. For total DNA extraction, 200 mg fecal samples were treated with QIAamp Fast DNA Stool Mini Kit (QIAGEN) following the manufacturer's protocol. Final DNA concentrations were quantified from the absorption at 260 nm with a Nanodrop 2000 spectrometer (Thermo Fischer Scientific).

2.5 16S rDNA sequencing

The V3-V4 variable region of 16S ribosomal RNA genes were amplified from fecal DNA extracts using universal primer set of Bakt 341F (5-cctacgggnggcwgcag) and Bakt 805R (5gactachygggtatctaatcc)³⁸. PCR amplification was performed with a 25 μ L reaction volume mixtures comprising 0.5 ng/µL of DNA template, 200 nM of each primer and 1x KAPA HiFi Hot Start Ready Mix (Kapa Biosystems) under the conditions: initial denaturation at 95°C for 3 min, 25 cycles of 95°C for 30 sec, 55°C for 30 sec and 72°C for 30 sec, followed by final extension at 72°C for 5 min. Amplification products were then purified using AMPure XP beads (Beckman Coulter). Subsequently, index PCR was performed using a 50 μ L reaction volume mixtures comprising 5 μ L of purified PCR products, 5 μ L of each index primer containing adapter sequence and sample specific 8 bp barcodes in the Nextera XT Index Kit v2 Set B (Illumina) and 1× KAPA HiFi Hot Start Ready Mix under the following conditions: 95°C for 3 min, 8 cycles of 95°C for 30 sec, 55°C for 30 sec and 72°C for 30 sec, followed by 72°C for 5 min. Resulting amplification products were purified using AMPure XP beads, quantified using the Qubit dsDNA HS Assay Kit (Invitrogen), and finally adjusted to 4 nM. Each amplicon was pooled and subjected to quantitative PCR using KAPA Library Quantification Kit Lightcycler 480 qPCR Mix (Kapa Biosystems), denatured following Illumina's guideline, and then adjusted to 4 pM. Amplicon library was combined with 5% of 4 pM PhiX Control v3 (Illumina) and subjected to pair-end sequencing using MiSeq instrument with a MiSeq 600-cycle v3 kit (Illumina). Resulting sequence reads were filtered for read quality, basecalled, and demultiplexed using bcl2fastq software (Illumina).

2.6 16S rDNA-based taxonomic analysis

Taxonomic analysis of FASTQ files produced by MiSeq was conducted by QIIME2 software (version 2019.7)³⁹. Quality-filtering, denoising, and removal of chimeric sequences were carried out by DADA2 plugin⁴⁰ using following parameters; --p-trim-left-f 17, --p-trim-left-r 21, --p-trunc-len-f 280, --p-trunc-len-r 200, --p-max-ee-f 2 –p-max-ee-r 2. Phylogenetic tree was constructed with FastTree⁴¹ after alignment with MAFFT⁴². Each feature was classified by a naïve-bayes classifier based on 99% sequence identity to the SILVA database (v.132). α -Diversity (observed OTUs and Shannon index) and β -diversity (weighted UniFrac distance) were calculated by Qiime2 pipeline. Statistical significance of β -diversity was determined by PERMANOVA test in Qiime2 pipeline.

2.7 Measurement of serum zonulin

Serum zonulin levels were determined by Mouse Haptoglobin ELISA Kit (Abcam, ab157714). The assay was performed according to the manufacture's recommended methods.

2.8 Bacterial translocation

The spleen was removed and immediately placed into 500 μ L of sterile Luria-Bertani (LB) medium too quantitate and identify bacteria. The spleen then was homogenized with a BioMasher (Nippi), and 200 μ L was plated on LB agar plates and cultured either under aerobic conditions for 24 h or anaerobic conditions for 48 h at 37°C. Colony-forming unit (CFU) were counted and calculated per organ.

2.9 Quantification of fecal Crps

Fecal samples were dried and powdered by using a Multi-beads shocker (Yasui Kikai). 30 mg of fecal samples was vortex mixed in 300 μ L PBS for 12 h at 4°C. The fecal suspension was centrifuged at 20,400 g for 10 min at 4°C, and Crp1 or Crp4 levels in supernatants were measured by sandwich ELISA as previously described^{34,43}.

2.10 Whole-mount immunofluorescent staining and image analysis

Whole-mount immunofluorescent staining was conducted using a modification of a previously reported method⁴⁴. The ileal tissue was fixed in 4% paraformaldehyde (Sigma) for 2 h at room temperature. Fixed tissue was permeabilized with 0.5 % Triton X-100 (Sigma) overnight at room temperature, and then blocked with 10% goat serum (Sigma) and 0.5% Triton X-100 overnight at 4°C. For SD and CDAHFD group, antibody reaction was performed using FITC labeled mouse anti-Crp1 antibody (50 µg/mL, clone 77-R63, produced in our laboratory) and Alexa Fluor 647-labeled antimouse/human CD324 (E-cadherin) antibody (1:100, clone DECMA-1, BioLegend). For CDAHFD+PBS and CDAHFD+R-Spo1 group, the primary antibody reaction was performed with rabbit anti-Olfactomedin 4 (Olfm4) antibody (1:80, clone D6Y5A, Cell Signaling) for 1 day at 4°C, and then the secondary antibody reaction was performed with Alexa Fluor 555 conjugated F(ab')2goat anti-rabbit IgG (dilution 1:500, Thermo Fisher Scientific) and FITC labeled mouse anti-Crp1 antibody overnight at 4°C. Following washing, nuclei were stained with DAPI (Thermo Fisher Scientific). Samples were immersed in the optical-clearing solution (RapiClear 1.52, Sunjin Lab). For quantification of Crp1 fluorescence intensity and counting numbers of Paneth cells and stem cells, Z-stack images were obtained using a confocal microscope (A1, Nikon) equipped with CFI Apo LWD $20X \text{ WI } \lambda S$ (Nikon). The number of Paneth cells was quantified by counting Crp1 immunostaining positive cells on 3 fields ($150 \times 150 \ \mu m^2$) per tissue. The number of stem cells was quantified counting Olfm4 positive cells on 3 fields ($150 \times 150 \ \mu m^2$) per tissue. The region of interest was created using image analysis software, NIS-Elements AR ver. 5.11 (Nikon), and the Crp1 fluorescence intensity per Paneth cell was measured on 3 fields ($150 \times 150 \ \mu m^2$) per tissue, and the mean intensity per field was calculated. For quantification of the number and diameter of Paneth cell granules, Z-stack images were obtained using A1 with CFI Apo TIRF 60X Oil (Nikon). The number and diameter of Paneth cell granules are diameter of Paneth cells/field) per tissue.

2.11 Transmission electron microscopy

5-mm-long segments of terminal ileum were immediately fixed in 2% paraformaldehyde and 2% glutaraldehyde at 4°C for overnight. Then, samples were post-fixed with 2% osmium tetroxide at 4°C for 2 h. Dehydration was carried out, followed by embedding in Quetol-812 epoxy resin (Nisshin EM). After staining with 2% uranyl acetate and lead stain solution (Sigma), the ultrathin sections were examined with a JEM-1400Plus transmission electron microscope (JEOL Ltd.) at an acceleration voltage of 100 kV.

2.12 Paneth cell RNA-seq analysis

Crypts were isolated from the small intestine of mice fed with SD or CDAHFD for 3 wk as previously described²⁷. In brief, the ileum segments were shaken in cold HBSS containing 30 mM EDTA. After vigorous shaking for ~300 times in HBSS, the isolated crypts were resuspended in HBSS containing 300 U/mL collagenase (Sigma), 10 mM Y-27632 (Sigma), and 1 mM N-acetylcysteine (Sigma), and shaken at 180 rpm for 5 min at 37 °C on a horizontal shaker (TAITEC). Then, 50 μ g/ μ L Dnase I (Roche) was added, and the sample was mixed by pipetting. Cells were pelleted at 500 g for 5 min at 4°C and resuspended in washing buffer (DMEM/F12 containing 10 μ M Y-27632 and 1 mM N-acetylcysteine), then passed through 40- μ m cell strainer (BD Falcon). Paneth cells were stained with

Zinpyr-1 (Santa Cruz) and Allophycocyanin (APC) anti-CD24 (clone M1/69,)Abcam) in washing buffer for 10 min at 37°C. Following Paneth cell labeling, the cells were sorted by flow cytometry using a cell sorter (JSAN, Bay Bioscience). Single cells were gated by forward scatter and side scatter. Cells were sorted directly into lysis buffer for RNA isolation (PureLink RNA Mini Kit, Invitrogen). To make a pooled sample, at least 10,000 Paneth cells were sorted from 2 separate mice, and sorting experiment was repeated three times. 2 pools (each pool contains Paneth cells from 6 separate mice) were prepared for each group. Total RNA was isolated by Invitrogen® PureLink RNA Purification System according to manufacturer's instructions, and cDNA was synthesized using a SMART-Seq. Sequencing libraries were built with the TruSeq RNA Library Prep Kit (Illumina) and then submitted to Illumina NovaSeq 6000 for 100-bp PE reads sequencing. Fragments per kilobase of transcript per million mapped reads (FPKM) values were used, genes with FPKM values below 1 were not included in the analysis, and fold change ≥ 1.5 or ≤ 0.67 was considered differentially expressed.

2.13 Visualization and quantification of Paneth cell granule secretion

Paneth cell granule secretion was assessed by a modification of a previously reported method²⁷. Isolated crypts from the distal 20 cm of small intestine of mice fed with SD and CDAHFD for 3 wk were embedded in Matrigel (Corning) on a collagen-coated 8 well chamber cover (Matsunami). After Matrigel polymerization, the enteroid culture media were added and incubated for 1 h at 37°C, 5% CO2. The enteroids were stimulated by 1 µM of carbamyl choline (CCh) (Sigma). The differential interference contrast images of Paneth cells before and 10 min after adding CCh were obtained from confocal microscopy (A1, Nikon). For quantification of Paneth cell granule secretion, area of the granules at pre-and post-stimulation was measured by using the image analysis software, NIS-Elements AR, and calculated percent granule secretion. Paneth cells in 10 crypts of each mouse (4 mice of each group) were evaluated.

2.14 Western blot analysis

Quantification of reduced-form Crp1 was performed as described previously⁴⁵. In brief, 10 mg of freeze-dried powdered fecal samples were suspended with 100 µl of 30% (vol/vol) acetic acid and vortexed at 4°C overnight. Fecal suspension was centrifuged at 15,000 g for 30 min at 4°C. Supernatants were concentrated in a SpeedVac concentrator (Thermo Savant) and resuspended in 20 µl of PBS. 1 µL of resuspension was reacted with 9 µL of 20 ng/µL trypsin solution, and samples were subjected to Tris-Tricine SDS–PAGE. Proteins were transferred to polyvinylidene fluoride membranes. The membranes were blocked by BSA-free StabilGuard Immunoassay Stabilizer (Surmodics) and incubated with anti-Crp1 antibody (1 mg/mL, clone 76-R29, produced in our laboratory). The blots were visualized with Chemi-Lumi One ultra (Nacalai Tesque). For quantification of Crps, band intensities were determined by ImageJ software.

2.15 Administration of R-Spo1 and Crp4

Recombinant human R-Spo1 provided from Kyowa Kirin Co., Ltd was generated as previously reported^{46,47}. R-Spo1 was intravenously administered at a dose of 600 µg three times a week for 3 wk. Recombinant Crp4, produced and purified as previously described⁴⁸, was orally administered at a dose of 110 µg twice daily for 6 wk.

2.16 Data availability

Paneth cell RNA-seq data were deposited in the NCBI's Sequence Read Archive (SRA PRJNA747067). 16S rRNA sequences were uploaded to the NCBI's SRA (PRJNA747068).

2.17 Statistical analyses

Data were analyzed with GraphPad Prism 8 software (Graphpad software), and results were represented as individual points with mean values and error bars representing SEM. Statistical significance between 2 groups was determined by unpaired two-tailed Student's t test. Pearson's test was used to analyze the correlation. A P value of less than 0.05 was considered statistically significant.

Chapter 3. Results

3.1 Dysbiosis occurs before onset of NASH in CDAHFD-fed mice

Histological analyses of liver were conducted on mice fed standard diet (SD) or CDAHFD. At 1 wk, fat accumulation and inflammatory foci were observed in the CDAHFD group, also both the area of steatosis and the number of inflammatory foci increased with the number of wks on CDAHFD (Fig. 1, a and b). Hepatocellular ballooning increased at 3 wk in the CDAHFD group and continued to increase until 12 wk. Sirius red staining was performed to evaluate hepatic fibrosis, showing that Sirius red-positive fibrotic areas first appeared at 3 wk and continued to increase until 12 wk (Fig. 1, a and b). The number of TUNEL-positive liver cells in the CDAHFD group increased at 3 wk and continued to increase with NASH progression up to the 12 wk endpoint (Fig. 1c). These results showed that mice fed with CDAHFD developed progressive hepatic fibrosis with steatosis, lobular inflammation, and apoptosis, which reproduces the typical disease development of human NASH⁹.



Figure 1. Hepatic steatosis, lobular inflammation, hepatocellular ballooning, and fibrosis occur

in the CDAHFD group.

(a) Representative images of H&E- and Sirius red-stained liver sections. Arrowheads indicate Sirius red-positive area. Scale bars: 50 μ m. (b) Quantification of steatosis area, the number of inflammatory foci and Sirius red-positive area. (c) Representative images of TUNEL staining of liver sections (left). Arrowheads indicate TUNEL-positive cells. Scale bars: 10 μ m. Quantification of TUNEL-positive cells (right). Data are shown as mean ± SEM for *n* = 3–6 per group. **P* < 0.05, ***P* < 0.01 and ****P* < 0.001, by unpaired 2-tailed Student's t test.

To examine whether endoplasmic reticulum (ER) stress, oxidative stress, and impaired autophagy, with known involvement in NASH pathology⁷ occur in the CDAHFD group, specific marker gene expression was quantified by real-time PCR. Among ER stress markers, *Pdia3* mRNA levels decreased at 1 wk and increased at 3 wk, *Chop* mRNA levels increased at 6 and 12 wk, and *Perk* mRNA was elevated at 12 wk with CDAHFD feeding compared to the SD group (Fig. 2a). Expression of *Nox2* mRNA, a key source of redox radicals, increased, and levels of antioxidant enzyme *Prdx6* mRNA decreased, both after 1 wk of CDAHFD feeding (Fig. 2b). Among autophagy activating protein genes, *Atg3* mRNA declined after 1 wk on CDAHFD followed by decreasing levels of both *Atg12* and *Lc3b* mRNAs after 3 wk of CDAHFD feeding (Fig. 2c). In addition, *F4/80* and *Cd11b*, markers of resident and invasive macrophages, respectively, and dendritic cell marker *Cd11c* mRNA levels continued to increase after 1 wk (Fig. 2d). Fibrosis related growth factor *Tgfb1* continued to increase after 3 wk of CDAHFD group (Fig. 2e). These results showed that inflammation and apoptosis in the liver contributed to the onset and progression of fibrosis and that ER stress, oxidative stress, and impairment of autophagy were involved in NASH development of the CDAHFD group.



Figure 2. The CDAHFD group develops NASH pathology associated with dysregulated expression of ER stress, oxidative stress, and autophagy related genes.

(a–e) Hepatic mRNA expression of ER stress (a), oxidative stress (b), autophagy (c), inflammation (d), and apoptosis related genes (e) was analyzed by real-time PCR. Data are shown as mean \pm SEM for *n* = 3–6 per group. **P* < 0.05, ***P* < 0.01 and ****P* < 0.001, by unpaired 2-tailed Student's t test.

To investigate whether dysbiosis occurs in the CDAHFD group, we conducted 16S rRNA gene sequencing on fecal DNA. α-Diversity analysis showed that observed both operational taxonomic units (OTUs) and Shannon index were significantly decreased in the CDAHFD group from 1 wk (Fig. 3a). Principal coordinates analysis (PCoA) showed that the composition of intestinal microbiota in the CDAHFD group was different from that of the SD group after 1 wk and continued through 12 wk (Fig. 3b). At the family level, the relative abundances of Eggerthellaceae, Bacteroidaceae, Prevotellaceae, Streptococcaceae, Peptostreptococcaceae, and Ruminococcaceae were significantly increased in the CDAHFD group. In contrast, Muribaculaceae, unassigned family of unassigned Bacteroidia class,

unassigned Clostridiales, Clostridiales VadinBB60 group, and Erysipelotrichaceae were significantly decreased relative to the SD group (Fig. 3c). At the genus level, a total of 18 genera showed significantly increased after CDAHFD feeding while total of 10 genera decreased in the CDAHFD group during disease progression (Fig. 3d). These results showed that dysbiosis occurs before the onset of NASH in mice fed CDAHFD.



Figure 3. Dysbiosis occurs before onset of NASH in the CDAHFD group.

(a) Observed OTUs and Shannon index in SD and CDAHFD groups. (b) PCoA plot of intestinal microbiota based on weighted UniFrac distance of SD and CDAHFD groups. (c, d) Relative abundance of individual genera that were significantly different between the SD and CDAHFD groups at the family (c) and genus level (d). Data are shown as mean \pm SEM for n = 5-6 per group. The data represent mean \pm SEM. *P < 0.05, **P < 0.01 and ***P < 0.001, by unpaired 2-tailed Student's t test.

Because increased intestinal permeability associated with dysbiosis contributes to NASH progression via the influx of intestinal bacteria and bacterial components into the liver, we next measured serum zonulin, a marker of intestinal permeability⁴⁹. At 1 wk, the SD and CDAHFD groups had similar serum zonulin levels. However, at 3 wk and continuing for 12 wk, serum zonulin levels in the CDAHFD group were increased significantly compared to the SD group sera (Fig. 4a). We further investigated whether bacterial translocation occurred. Bacterial colonies were detected in the spleen at 6 wk in the CDAHFD group (Fig. 4b). These results indicated that intestinal hyperpermeability accompanied by dysbiosis occurred in the CDAHFD group.



Figure 4. Increased intestinal permeability and bacterial translocation to spleen occurs in the CDAHFD group.

(a) Serum zonulin levels in SD and CDAHFD group. (b) Quantification of bacteria in spleen by CFUs cultured from spleen of SD and CDAHFD group at 3, 6, and 12 wk under anaerobic condition. Data are shown as mean \pm SEM for n = 3-6 per group. The data represent mean \pm SEM. *P < 0.05, **P < 0.01 and ***P < 0.001, by unpaired 2-tailed Student's t test

3.2 Paneth cell α -defensin secretion decreases and relates to NASH dysbiosis

Because Paneth cell α -defensin have a role as regulator of the intestinal microbiota, we next examined whether Paneth cell α -defensins are involved in the onset of NASH in the CDAHFA group. To evaluate the quantity of α -defensins secreted by Paneth cells, fecal levels of Crp1, one of the most abundant Crps in the small intestine⁵⁰, were measured. Before the onset of liver fibrosis, Crp1 secretion in the CDAHFD group significantly decreased compared to the SD group and continued for 12 wk (Fig. 5).



Figure 5. Secretion of Paneth cell α-defensin decrease in the CDAHFD group.

Quantification of fecal levels of Crp1. Data are shown as mean \pm SEM for n = 5-6 mice/group. *P <

0.05 and **P < 0.01, by unpaired 2-tailed Student's t test.

To determine whether decreased α -defensin secretion is associated with dysbiosis and NASH pathogenesis, correlation analyses among quantity of α -defensin levels in feces, the abundance of the intestinal microbiota, and NASH pathology were conducted. Among 28 genera at the genus level, which showed significantly different proportion between the SD and CDAHFD groups during disease progression (Fig. 3d), twenty genera showed a positive or negative correlation with fecal Crp1 (Fig. 6). Among them, the relative abundance of 10 genera which increased in the CDAHFD group, including Bacteroides, Alloprevotella, GCA-900066575, Lachnoclostridium, uncultured Lachnospiraceae, unassigned Peptostreptococcaceae, Harryflintia, Ruminiclostridium 5. Ruminiclostridium 9, and UCG-009, correlated positively with NASH pathology including steatosis, inflammation, apoptosis, and fibrosis. On the other hand, the 6 genera that decreased in abundance, i.e., Muribaculum, unassigned Muribacululaceae, Muribaculaceae uncultured bacterium, NK4A136 group, unassigned genus of the unassigned Clostridiales family, and uncultured Clostridiales VadinBB60 group, correlated negatively (Fig. 6). Our results identified total 16 genera which related with both Crp1 reduction and NASH pathophysiology, suggesting that Crp1 reduction contributed to NASH progression via alterations in these intestinal bacteria.



Figure 6. The CDAHFD group shows dysbiosis correlated with both the quantity of fecal αdefensin and NASH pathology.

28 genera that were significantly increased or decreased in CDAHFD group compared with SD group. Red indicates genera significantly increased in CDAHFD group, and blue indicates genera significantly decreased in CDAHFD group. Heatmap showing Pearson correlation coefficients between relative abundance of significantly changed genera in CDAHFD group and fecal Crp1 levels from Fig. 5 or NASH pathology from Fig. 1b. Correlation analysis between Crp1 levels and relative abundance of individual genera was conducted using the data of 1, 3, 6, and 12 wk. Correlation analysis between NASH pathology and relative abundance of individual genera was conducted using the data of 12 wk. $*P \le 0.05$, $**P \le 0.01$ and $***P \le 0.001$, by Pearson correlation test.

To investigate the cause of decreased Crp1 secretion, Crp1 mRNA expression, Paneth cell number, and whole-mount fluorescence immunostaining of Crp1 were performed on the ileal tissue samples. For both Crp1 mRNA expression and Paneth cell number, no differences were observed between the SD and CDAHFD groups (Fig. 7, a–d). In contrast, fluorescent intensities of Crp1 in the CDAHFD group were diminished compared to the SD group after 3 wk (Fig. 7, c and e), indicating protein expression of α -defensin decreased in Paneth cells of the CDAHFD group.

Because α -defensins are abundant in Paneth cell granules under normal conditions, we examined the morphology of Paneth cell granules to possibly explain the decreased levels of α -defensins in Paneth cells of the CDAHFD group. The diameter of Paneth cell granules decreased after 1 wk of the CDAHFD group, and the numbers of Paneth cell granules also decreased from 3 to 12 wk (Fig. 7, f– h). Transmission electron microscopy images confirmed the abnormal morphologies of Paneth cell granules in the CDAHFD group (Fig. 7i).



Figure 7. Paneth cells of the CDAHFD group showed decreased protein expression of α -defensin and abnormal granules.

(a, b) mRNA levels of Crp1 in ileum of SD and CDAHFD group at 3 wk (a), and 6 wk (b) analyzed by real time-PCR. Data are shown as mean \pm SEM for n = 3 mice/group. (c) Representative images of immunofluorescence staining of Crp1 (green) in small intestine from SD and CDAHFD group. DAPI (blue) stains the nucleus. Scale bars: 20 µm. (d, e) Quantification of the number of Paneth cells (d) and Crp1 fluorescence intensity on Paneth cells (e) shown in c. Data are shown as mean \pm SEM for n

= 9–18 fields/group and quantified based on 3 fields per mouse. (f) Representative images of whole mount small intestine obtained with confocal microscopy. Scale bar: 5 μ m. (g, h) Granule diameter (g) and granule number (h) of each Paneth cell per field shown in f. Data are shown as mean ± SEM for *n* = 9–18 fields/group and quantified based on at least 3 fields per mouse. Each group contains 3– 6 mice. (i) Representative images of transmission electron microscopy of Paneth cells at the base of the ileal crypts in SD and CDAHFD group at 1, 3, 6, and 12 wk. Paneth cells in SD group have tightly packed electron-dense core granules, in contrast, Paneth cells in CDAHFD group have fewer loosely packed granules with a halo appearance. *n* = 1 for each group. Scale bars: 5 μ m. (a, b, d, e, g, h) **P* < 0.05, ***P* < 0.01 and ****P* < 0.001, by unpaired 2-tailed Student's t test. NS, not significant.

To further determine the molecular basis for these functional alterations in Paneth cells, RNA sequencing (RNA-seq) analysis was performed on isolated small intestinal Paneth cells from the SD and CDAHFD groups. One hundred and forty-five genes were upregulated in CDAHFD mice compared to the SD group, while 2,202 genes were downregulated in the CDAHFD group (Fig. 8a). These differentially regulated genes were compared with gene products by referring to GO terms: Secretion (GO: 0046903) and trans-Golgi network (GO: 0005802), known to be important in the granule secretory pathway. Twelve upregulated genes in CDAHFD Paneth cells include ion transporter *Car4*, oxidative stress gene *Chac1*, glycosylation protein *Tmem165*, and secretion protein *Stxbp1*. The 97 down regulated genes include ion transporters *Cacr2a*, *Kcnq1*, *Cftr*, and *Kcnn4*, vesicle transporters *Optn*, *Trappc9*, *Vamp2*, *Dop1a*, *Rab26*, *Lrba*, *Arfrp1*, and *Gga3*, autophagy related proteins *Klhl20*, *Nlrp6*, *Ap4m1*, and *Bsn*, and oxidative stress related protein *Arntl1* (Fig. 8b). These results indicated that expression of genes related to the granule secretory pathway, including vesicle transporters and ion channel transporters, is dysregulated in Paneth cells of the CDAHFD group. Thus, to evaluate

Paneth cell granule secretion of the CDAHFD group directly, granule secretory responses of Paneth cells were visualized in enteroids, a three-dimensional culture system of small intestinal epithelial cells²⁷. When CCh, which induces Paneth cell granule secretion, was added to enteroid cultures derived from small intestinal crypts of the CDAHFD group, the quantity of granules secreted by Paneth cells was significantly lower than that of SD group enteroids (Fig. 8c). These results revealed that Paneth cells in the CDAHFD group decreased α -defensin secretion into the intestinal lumen due to decreased protein expression of α -defensin and impaired granule secretory function.





(a) Scatterplot of global gene expression profiles of SD and CDAHFD group derived from RNA-seq analysis. (b) Heatmap showing log2 fold change of differentially expressed genes (1.5-fold increase or decrease) associated with Paneth cell functions in CDAHFD group compared with SD group. (c) Enteroids derived from the small intestine of SD and CDAHFD groups were stimulated by 1 mM CCh for 10 min. Percent granule secretion was calculated as percent area granule secretion. Data are shown as mean \pm SEM for n = 40 crypts/group. Each group contains 4 mice. *P < 0.05, by unpaired 2-tailed Student's t test.

It has been reported that abnormal Paneth cells secrete reduced-form α -defensin that lack disulfide bonds resulting in disruption the spectrum of bactericidal activity and are involved in the development of ileitis via dysbiosis in Crohn's disease model mice⁴⁵. Therefore, we further measured the amount of reduced-form Crp1 (rCrp1) in feces of the CDAHFD group. Oxidized-form crp1 (oxCrp1) decreased in the CDAHFD group compared to the SD group, whereas rCrp1 was not detected in both at 3 and 12 wk (Fig. 9, a and b). These results indicated that Paneth cell dysfunction impairs granule secretion, resulting in diminished release of native, i.e., oxidized-form, α -defensins into the intestinal lumen.



Figure 9. Paneth cells in the CDAHFD group do not secrete reduced-form α-defensins.

Tricine SDS-PAGE Western blot analysis of oxCrp1 or rCrp1 in fecal extracts of SD and CDAHFD

group at 3 wk (a) and 12 wk (b). Chemically synthesized ox and rCrp1 were used as positive controls. Each sample was treated with trypsin (Trp+) or PBS (Trp–). Crps detected in Trp– were considered to be the sum of ox and rCrp1, and Crps detected in Trp+ were considered as rCrp1 that escaped tryptic digestion. The amount of ox and rCrp1 in fecal extracts. The amount of oxCrp1 in each sample was calculated from band intensity of Trp+, and the amount of rCrp1 was calculated from the difference between band intensity of Trp– and Trp+. Data are shown as mean ± SEM for n = 5-6 per group. The data represent mean ± SEM. *P < 0.05 and **P < 0.01 by unpaired 2-tailed Student's t test. NS, not significant.

3.3 Restoration of α -defensin by R-Spo1 administration suppresses dysbiosis and prevents progression of NASH

To clarify the association between the early reduction of secreted luminal Paneth cell α -defensins and dysbiosis along with NASH progression, we tested whether restoration of α -defensins prevented dysbiosis induced by CDAHFD feeding. Because Wnt activator R-Spo1 enhances α -defensin secretion by stimulating enhanced Paneth cell differentiation from stem cells³⁵, CDAHFD-fed mice were injected intravenously with R-Spo1 for 3 wk. Measurements of fecal Crp1 tested whether R-Spo1 treatment restored luminal levels of secreted Crp1. Fecal Crp1 levels in the R-Spo1-treated CDAHFD mice (CDAHFD+R-Spo1 group) were increased at 1 wk, and after 2 wk, significantly increased compared to CDAHFD-fed mice treated with PBS (CDAHFD+PBS group) (Fig. 10a). Paneth cell numbers in CDAHFD+R-Spo1 mice were increased significantly at 3 wk along with stem cells, showing that R-Spo1 treatment restored luminal Crp1 levels by elevating Paneth cell numbers (Fig. 10b).

Next, we assessed whether enhanced luminal Crp1 levels by R-Spo1 treatment attenuates dysbiosis. Among the families of intestinal bacteria which increased in mice fed CDAHFD during disease progression (Fig. 3c), the relative abundances of Prevotellaceae and Peptostreptococcaceae were reduced by R-Spo1 treatment. In contrast, R-Spo1 treatment decreased the proportion of enteric microbiota that showed increases during disease progression, including Muribaculaceae, unassigned Clostridiales, and unassigned family of unassigned Bacteroidia (Fig. 10c). Fecal Crp1 levels were positively correlated with the relative abundance of Muribaculacea and unassigned Clostridiales (Fig. 10c). At the genus level, among the intestinal bacteria that correlated with both fecal Crp1 levels and NASH pathology (Fig. 3d), the proportion of *Muribaculum*, unassigned Muribaculaceae, Muribaculaceae uncultured bacterium, and unassigned Clostridiales was restored by R-Spo1 treatment, showing positive correlation with Crp1 levels. Also, the outgrowth of *Harryflintia* was inhibited, showing negative correlation with Crp1 levels (Fig. 10d). These results indicated restoration of luminal α -defensin by R-Spo1 administration suppressed dysbiosis related to NASH.



Figure 10. R-Spo1 restores luminal α-defensin and suppresses dysbiosis.

Six-week-old C57BL/6J mice were fed CDAHFD to induce NASH and intravenously injected with R-Spo1 at a dose of 600 µg or PBS three times a week for 3 wk. (a) Fecal Crp1 levels in the CDAHFD+PBS and CDAHFD+R-Spo1 groups. (b) Representative images of immunofluorescence staining of Crp1 (green) and Olfm4 (red) in small intestine from CDAHFD+PBS and CDAHFD+R-Spo1 mice at 3 wk (left). Arrowheads indicate Paneth cells and asterisk indicate stem cells. Scale bar:

20 µm. Quantification of the number of Paneth cells assessed by expression of Crp1 and the number of stem cells assessed by expression of Olfm4 per ileal unit area (right). Data are shown as mean \pm SEM for *n* = 6 fields/group and quantified based on at least 2 fields per mouse. Each group contains 3 mice. (c) Relative abundance of individual genera that were significantly recovered in CDAHFD+R-Spo1 group from CDAHFD mice of 11 family in Fig. 3c (upper panel). Correlation analysis between fecal Crp1 levels and relative abundance of individual genera CDAHFD+PBS and CDAHFD+R-Spo1 group (lower panel). (d)Relative abundance of individual genera that were significantly recovered in CDAHFD+R-Spo1 group from CDAHFD group of 28 genera shown in Fig. 3d (upper panel). Correlation analysis between fecal Crp1 levels and relative abundance of individual genera (lower panel). *r* and *P* values by Pearson correlation test are presented. Data are shown as mean \pm SEM for *n* = 3 per group. **P* < 0.05, ***P* < 0.01 and ****P* < 0.001, by unpaired 2-tailed Student's t test.

To discern the suppression of dysbiosis due to restoration of α -defensin by R-Spo1 administration prevented NASH development, histological analysis of liver was conducted. The area of steatosis, the numbers of inflammatory foci, hepatocellular ballooning, and the area of fibrosis all were decreased significantly in the CDAHFD+R-Spo1 group compared to the CDAHFD+PBS group (Figure 11, a and b). R-Spo1 treatment significantly decreased hepatic mRNA levels of *Nox2*, *F4/80*, *Tgfb1*, *Trailr2*, and *Bax*, tended to decrease *Cd11c* expression, and decreased the number of TUNEL-positive cells significantly (Fig. 11, c and d). These results indicated that R-Spo1 treatment attenuated steatosis, inflammation, apoptosis, and fibrosis in the livers of mice fed CDAHFD. Among the genus of intestinal bacteria corrected for abnormal occupancy by R-Spo1 treatment (Fig. 10d), both *Muribaculum* and unassigned Muribaculaceae showed negative correlation with the steatosis area, the number of TUNEL-positive cells, and the area of fibrosis, and unassigned genus of the unassigned Clostridiales family was negatively correlated with numbers of TUNEL-positive cells (Fig. 11e). The number of inflammatory foci was negatively correlated with Muribaculaceae uncultured bacterium, positively correlated with *Harryflintia*, and tended to be negatively correlated with *Muribaculum*, Muribaculaceae uncultured bacterium and unassigned genus of the unassigned Clostridiales family (Fig. 11e). In addition, fecal Crp1 levels were negatively correlated with the number of inflammatory foci, the number of TUNEL-positive cells, and the fibrosis area, indicating restoration of Crp1 secretion ameliorated NASH pathology (Fig. 11f). Taken together, these results indicated that enhancing luminal α -defensin secretion from Paneth cells by R-Spo1 treatment reduces liver fibrosis via ameliorating inflammation and apoptosis in the liver along with suppressing dysbiosis.



Figure 11. Restoration of α -defensin by R-Spo1 administration improves NASH progression by suppressing dysbiosis.

(a) Representative images of H&E- and Sirius red-stained liver sections. Scale bars: 50 μm. (b)
Quantification of steatosis area, the number of inflammatory foci and Sirius red-positive area. (c)
Hepatic mRNA expression of ER stress, oxidative stress, autophagy, inflammation, and apoptosis
marker genes. (d) Representative images of TUNEL staining of liver sections (upper). Scale bars: 10

 μ m. Quantification of TUNEL-positive cells (lower). (e) Correlation analysis between relative abundance of individual genera and NASH pathology. (f) Correlation analysis between fecal Crp1 levels and NASH pathology. Data are shown as mean ± SEM for *n* = 3 per group. **P* < 0.05 and **P < 0.01 by unpaired 2-tailed Student's t test (b–d) and Pearson correlation test (e and f).

3.4 Oral administration of α -defensin suppresses dysbiosis and ameliorates NASH pathology

Finally, since granules of Paneth cells contain microbicidal components in addition to abundant α defensins⁵¹, we investigated whether oral administration of an exogenous α -defensin prevents dysbiosis and subsequent NASH development. Crp4 has the most potent in vitro bactericidal activities among Crps⁵², and oral administration of Crp4 improved the homeostasis of the intestinal microbiota in GVHD model and in a chronic social defeat stress model^{35,53}. Therefore, Crp4 was administered orally to CDAHFD-fed mice for 6 wk. Because C57BL/6 strain lacks the Crp4 gene⁵⁴ and dosing with Crp4 introduces an exogenous α -defensin, we confirmed that orally administered Crp4 reached the intestinal lumen by measuring levels of fecal Crp4 (Fig. 12a). To assess whether oral Crp4 suppressed dysbiosis, the intestinal microbiota composition was analyzed. At the family level, among the intestinal bacteria that increased in the CDAHFD group during disease progression, the proportion of Eggerthellaceae, Streptococcaceae, and Ruminocccaceae decreased with Crp4 administration (Fig. 12b). At the genus level, the outgrowth of *Harryflintia* was inhibited by Crp4 administration comparably to R-Spo1 treatment. On the other hand, the increase of uncultured Lachnospiraceae and *Ruminoclostridium 9*, which were not reversed by R-Spo1 treatment decreased with Crp4 administration (Fig. 12c). These results indicated that oral administration of α -defensin suppressed dysbiosis related to NASH, correcting abnormal proportion of the intestinal bacteria partially different from R-Spo1 administration.



Figure 12. Oral administration of Crp4 suppresses dysbiosis.

Six-week-old C57BL6J mice were fed with CDAHFD to induce NASH and orally administered with Crp4 at a dose of 110 µg twice daily for 6 wk. (a) Fecal levels of Crp4 in CDAHFD+Crp4 group were analyzed by sandwich ELISA at 0 and 6 wk. (b) Relative abundance of individual genera at family level that were significantly recovered in CDAHFD+Crp4 group from CDAHFD group of 13 family in Fig. 3c. (c) Relative abundance of individual genera at genus level that were significantly recovered in CDAHFD group of 28 genera shown in Fig. 3d. Data are shown as mean \pm SEM for n = 6-8 per group. *P < 0.05, **P < 0.01 and ***P < 0.001, by unpaired 2-tailed Student's t test.

To assess whether oral administration of Crp4 ameliorated liver fibrosis by suppressing dysbiosis, histological analysis of liver was performed. The number of inflammatory foci, hepatocellular ballooning, and the area of fibrosis significantly decreased by oral administration of Crp4, even though the area of hepatic steatosis was not affected (Fig. 13, a and b). Analysis of liver gene expression revealed that Crp4 administration increased *Atg12* mRNA levels, decreased *Cd11b* and *Tgfb1* mRNAs, and tended to decrease levels of *Nox2* and *Cd11c* mRNAs. Genes that decreased in expression by R-Spo1 treatment including *F4/80*, *Trailr2*, and *Bax* were unaffected by oral Crp4 administration (Fig. 13c). TUNEL-positive cells in the liver decreased significantly in the Crp4-treated mice compared to untreated CDAHFD-fed mice (Fig. 13d). Moreover, both *Harryflintia* and *Ruminiclostridium 9*, which decreased by Crp4 administration, were positively correlated with the number of inflammatory foci and TUNEL-positive cells, showing that Crp4 ameliorated inflammation and apoptosis followed by fibrosis in the liver via suppressing dysbiosis (Fig. 13e).

Taken together, R-Spo1 and Crp4 prevented liver fibrosis with ameliorating inflammation and apoptosis via suppressing dysbiosis, though, the gene expression in the liver and the intestinal microbiota, which improved by Crp4 administration, were different from those by R-Spo1 treatment, suggesting different underlying mechanism.



Figure 13. Oral administration of α -defensin increases autophagy related gene expression and ameliorates liver fibrosis by suppressing dysbiosis.

(a) Representative images of H&E- and Sirius red-stained liver sections. Scale bars: 50 μ m. (b) Quantification of steatosis area, the number of inflammatory foci and Sirius red-positive area. (c) Hepatic mRNA expression of ER stress, oxidative stress, autophagy, inflammation, and apoptosis marker genes. (d) Representative images of TUNEL staining of liver sections (left). Scale bars: 10 μ m. Quantification of TUNEL-positive cells (right). (e) Correlation analysis between relative abundance of individual genera and NASH pathology. Data are shown as mean \pm SEM for n = 6-8 per group. **P* < 0.05, ***P* < 0.01 and ****P* < 0.001, by unpaired 2-tailed Student's t test (b, c, d) and Pearson correlation test (e).

Chapter 4. Discussion

4.1 Relationship between Paneth cell α-defensin and dysbiosis associated with NASH

NASH is one of the largest urgent unmet medical needs with continuously increasing numbers of patients worldwide. Recently, fibrosis was shown to be the major determinant of mortality in patients with NASH, so that understanding the mechanisms of fibrosis development in NASH is extremely important^{3,55,56}. This study focused on undetermined mechanisms of the "gut-liver axis" that lead to liver fibrosis in NASH pathology and analyzed the relationship between Paneth cell α -defensins and NASH pathology via dysbiosis. Although high-fat diet (HFD) induces NAFLD, HFD itself generally does not induce severe NASH pathology with fibrosis in mouse models⁵⁷. Therefore, we chose CDAHFD-induced mouse model of NASH, which have been reported to induce progressive liver fibrosis with acceleration of fat accumulation in the liver⁵⁸. In this study, CDAHFD model showed hepatic inflammation and apoptosis accompanied by ER stress, oxidative stress, and impairment autophagy in the liver during disease progression resemble to NASH patient⁷. Furthermore, we have clarified that the CDAHFD group exhibits dysbiosis, sharing some similar features with NAFLD/NASH patients. α-Diversity of the intestinal microbiota decreased in the CDAHFD group consistent with the previous report⁵⁹. At the family level, Prevotellaceae, reported to increase in NAFLD patients¹⁷, was elevated in the CDAHFD group. At the genus level, *Bacteroides*, known to increase in NASH patients and positively correlated with fibrosis score⁶⁰, was also elevated in the CDAHFD group. These results suggested that CDAHFD model is a suitable model for analyzing the relationship between NASH pathology and the intestinal homeostasis.

Dysbiosis has been reported to occur in NASH patients, but the cause of dysbiosis has not been well understood because the composition of the intestinal microbiota is affected by varied factors including diet, environmental factors, and host-derived factors⁶¹. In this study, Crp1, an abundant mouse α - defensin, significantly decreased before NASH onset along with disease progression and increased α defensin levels by R-Spo1 administration suppressed dysbiosis and ameliorated steatosis, inflammation, apoptosis, and fibrosis in the liver. R-Spo1 enhanced luminal secretion of α -defensin by regeneration of Paneth cells, though, Paneth cells secrete other antimicrobial proteins including lysozyme and secretory phospholipase A2⁶². In addition, R-Spo1 is reported to increase not only numbers of Paneth cells but also goblet cells³⁵ secreting mucus, which harbors the intestinal microbiota and influence its composition⁶³. Therefore, we further tested whether oral administration of α -defensin prevents NASH, and oral administration of Crp4 suppressed dysbiosis and attenuated NASH pathology. These results indicated that reduction of α -defensin caused dysbiosis and further promoted the development of NASH.

Because secretion of misfolded reduced-form α -defensin from abnormal Paneth cells has been reported to be related to dysbiosis in Crohn's disease model mice⁵³, whether rCrp1 is secreted in CDAHFD-fed mice was investigated. rCrp1 was not observed in the CDAHFD group, indicating that not abnormal quality but decreased quantity of α -defensin is involved.

4.2 Involvement of Paneth cell α -defensin in NASH pathology through regulation of the intestinal microbiota

While both R-Spo1 and Crp4 inhibited the outgrowth of *Harryflintia*, reported to be increased in the animal model of hyperlipidemia and atherosclerosis⁶⁴, they affected the intestinal microbiota differently, i.e., R-Spo1 restored Muribaculaceae, whereas Crp4 inhibited the outrgrowth of *Ruminiclostridium 9*. The composition and function of the intestinal microbiota differ between the mucus side and the luminal side in the intestine, forming a unique ecosystem⁶⁵. Muribaculaceae is known as the mucus-resident commensal bacteria⁶⁶, while *Ruminiclostridium 9* is reported to be a characteristic luminal bacteria⁶⁷. It is considered that secreted antimicrobial peptides including α -

defensin, do not easily diffuse into the intestinal lumen but are trapped in the mucus layer⁶⁸. Furthermore, human Paneth cell α -defensin HD5 does not elicit bactericidal activities against Muribaculaceae, suggesting a promotion of colonization³¹. Therefore, it is possible that intravenously administered R-Spo1 enhanced α -defensin secretion from *de novo* Paneth cells into mucus layer and predominantly contributed to modifying mucosa-associated bacteria including Muribaculaceae. In contrast to R-Spo1, orally administered α -defensin is expected to diffuse into the intestinal lumen. Thus, it is considered that orally administrated Crp4 mainly affected luminal resident bacteria including *Ruminiclostridium 9*. These findings provide an insight that R-Spo1 and Crp4 modified bacteria localized in mucus and lumen, respectively.

Although both R-Spo1 and Crp4 administration improved hepatic inflammation and fibrosis, R-Spo1 ameliorated steatosis and decreased apoptosis associated genes including *Trail2* and *Bax*, whereas Crp4 increased expression of Atg12, an autophagy related protein. Because various metabolites produced by intestinal bacteria, such as SCFA and bile acids, are known to be involved in NASH pathogenesis¹⁶, it is possible that each intestinal bacteria affects by different mechanisms.

Muribaculaceae, which increased by R-Spo1 treatment, is required for inner mucus layer formation of the intestine and SCFA production, known to have beneficial effects on the host^{69,70}. It has been reported that SCFA supplementation improved high fat diet-induced steatosis in liver with reduction of fatty acid transporter and improving lipid metabolism⁷¹. The expression of *Trailr2*, which decreased by R-Spo1 treatment, has been reported to be increased by free fatty acids in hepatocyte⁷². Therefore, it is suggested that R-Spo1 prevented steatosis, apoptosis, and fibrosis with increasing proportion of Muribacluaceae possibly accompanied by increasing SCFA production.

On the other hand, *Ruminiclostridium 9*, decreased by Crp4 administration in this study, increases in a high-fat/high-fructose diet-induced dyslipidemia model⁷³. In addition, *Ruminiclostridium 9* was positively correlated with the number of inflammatory foci in this study, suggesting *Ruminiclostridium*

9 is deleterious microbiota. *Atg12* whose gene expression increased by Crp4 has a crucial role for autophagy⁷⁴, and loss of autophagy in hepatocytes causes apoptosis, inflammation, and fibrosis in the liver⁷⁵. These findings suggested that oral administration of Crp4 prevented liver fibrosis by inhibition of apoptosis induced by impairment of autophagy and inflammation along with inhibiting bacterial outgrowth of *Ruminiclostridium* 9, which is a different mechanism of action from R-Spo1 treatment. Taken together, the different pathological features between R-Spo1 and Crp4 may be related to the difference in the improved intestinal microbiota composition, suggesting that α -defensin is involved in NASH pathology by regulating bacteria affecting liver function in both intestinal mucus and lumen.

4.3 Functional alterations of Paneth cells in NASH

Paneth cells regulate the composition of the intestinal microbiota by secreting granules rich in α defensins in response to cholinergic agents, bacteria, bacterial antigen, and certain dietary factors^{26,27,76}. Secretory responses of Paneth cells need a biphasic increase in cytosolic Ca²⁺ concentration, and Ca²⁺activated potassium channel KCNN4 is the essential modulator of Ca²⁺ concentration during Paneth cell secretion⁷⁷. Mice having defective cystic fibrosis transmembrane conductance regulator (CFTR) show unusual accumulation of Paneth cell granules in the intestinal crypt lumen, suggesting that CFTR is important for dissolution of secreted granules⁷⁸. In this study, the RNA-seq analysis of Paneth cells revealed that the expression of both *KCNN4* and *CFTR* decreased in Paneth cells of the CDAHFD group, suggesting that influx of ions such as Ca²⁺ is impaired. RAS-related GTP-binding protein (Rab) and soluble NSF attachment protein receptor (SNARE) family proteins, which allow vesicle transport and fusion, have key roles in granule secretion. The expression of *Rab26*, which has been reported to localize on secretory granules and required to amylase release in parotid acinar cells⁷⁹. Similarly, vesicle-associated membrane protein 2 (VAMP2) has an essential role for glucagon-like peptide 1 (GLP-1) secretion in intestinal L cell⁸⁰. Paneth cell in the CDAHFD group showed decreased expression of *Rab26* and *Vamp2*, which are essential for granule secretion. On the other hand, syntaxinbinding protein 1 (Stxbp1), whose overexpression inhibits the SNARE complex assembly and decreases insulin secretion⁸¹, increased in the CDAHFD group. Furthermore, decreased granule secretory function in the CDAHFD group was directly revealed by visualization and quantification of granule secretory response in enteroids. Because mutation of autophagy related genes such as *Atg16L1* leads to the abnormal morphology of Paneth cell granules, autophagy elicits important roles in granule formation and secretion of Paneth cells⁸². It has been reported that *Optn*, which decreased in the CDAHFD group Paneth cells, promotes autophagosome formation via recruitment of Atg12-5-16L1 complex⁸³. Our findings that Paneth cells in CDAHFD group showed reduction of *Optn* expression suggests that abrogation of autophagy leads to abnormal granule formation and decreased granule secretion. Taken together, it is possible that decreased granule secretion in mice fed CDAHFD occurs through impairment of influx of ions such as Ca²⁺, vesicle transport, and autophagy process in Paneth cells.

4.4 Potential prevention and treatment of NASH targeting Paneth cell α-defensin

There are currently no approved drugs to treat NASH, and the basis of all therapeutic interventions is lifestyle changes including diet and exercise². This study showed the possibility that decrease of Paneth cell α -defensin secretion could be the cause of onset and the progression of NASH via dysbiosis. The relationship between diet and Paneth cell function has been reported. Leucine, a dietary factor, and butyrate, generated by the metabolism of dietary fiber by the intestinal bacteria, induce the secretion of α -defensin from Paneth cell⁷⁶. N-3 polyunsaturated fatty acids increased expression of antimicrobial peptides, including α -defensin, and ameliorated intestine atrophy in total parental nutrition model mice⁸⁴. In addition, lactate increases expansion of Paneth cells by enhancing wnt signaling⁸⁵, which is expected to increase α -defensin secretion. These findings suggest that dietary

intervention targeting Paneth cell α -defensin could be a novel method of NASH prevention and treatment.

Although CDAHFD model mice has limitations such as not showing obesity and insulin resistance usually observed in patients with NASH⁸⁶, our findings provide novel insights into the process of NASH fibrosis in the "gut-liver axis" and possibly propose a novel therapeutic approach for NASH via regulation of the intestinal microbiota targeting Paneth cell α -defensin.

Chapter 5. Conclusion

This study revealed that decrease of Paneth α -defensin secretion associated with impairment of granule secretory function exacerbated liver apoptosis and inflammation via dysbiosis, contributing to development of NASH fibrosis. Both stimulating differentiation of Paneth cells to increase α -defensin secretion by R-Spo1 and oral administration of α -defensin improved dysbiosis and prevented fibrosis. Taken together, our findings provide novel insights into NASH pathophysiology in the "gut-liver axis" and further propose a novel therapeutic approach for NASH targeting Paneth cell α -defensin via regulation of the intestinal microbiota.

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