The gut microbiome in patients with chronic lymphocytic leukemia


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The gut microbiome in patients with chronic lymphocytic leukemia

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Authors’ Contribution
CUN, RS, and TF designed the study, EI, CC and CUN were responsible for sample collection and inclusion of patients, MNJ and RP performed the sequencing, TF, MJ, and CRM performed the bioinformatic and statistical analyses, TF and CUN wrote the first version of the manuscript, all authors read, contributed and approved the final version of the manuscript.

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CUN received research funding and/or consultancy fees from Abbvie, AstraZeneca, CSL Behring, Janssen, Octapharma, Roche and Takeda outside this work. MNJ is founder and shareholder of Nano1Health S.L.
Data sharing statement
The datasets generated and analyzed during this study are derived from patients treated in Denmark. The datasets contain sensitive patient data governed by GDPR and Danish law. Due to Danish legislation (Act No. 502 of 23 May 2018) and approvals granted by the Danish Data Protection Agency, it is not possible to upload raw data to a publicly available database. However, access to these data can be made available from the corresponding author on reasonable request, provided a data transfer agreement is entered into according to current regulations.

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The gut microbiome, an ecosystem formed by commensal, symbiotic, and pathogenic microorganisms colonizing the gastrointestinal tract, may impact both immune function and carcinogenesis. The host immune system plays a vital role in the maintenance of gut microbiome homeostasis by establishing a balance between eliminating invading pathogens and promoting the growth of beneficial microbes. When this balance is disturbed, a state of dysbiosis arises in the microbial ecosystem. Conditions found to be associated with gut dysbiosis include, but are not limited to, inflammatory disorders including inflammatory bowel diseases (IBD), diabetes, as well as obesity, and asthma. The gut microbiome also seems to influence cancer susceptibility, and to correlate with tumorigenesis and progression. Within the area of hematological malignancies, studies focusing on acute leukemias, lymphoproliferative disorders, and multiple myeloma, have found microbiome dysbiosis and decreased microbiome diversity to be related to microenvironmental alterations induced by the disease itself, chemotherapy, and/or antibiotics.

To our knowledge, no previous studies have investigated the gut microbiome in patients with chronic lymphocytic leukemia (CLL). As CLL represents an antigen driven malignancy with immune dysfunction, the gut microbiome could both be implicated in the pathogenesis of CLL through antigenic drive, and contribute to the distortion of the immune system. However, the CLL microbiome itself may also be impacted by the immune dysfunction as well as reflect the increased prescription of antimicrobials for this patient group. Thus, our study focusing on the fecal microbiome in patients with CLL aimed to describe perturbations in the gut microbial composition, and to characterize potential signature for CLL-related gut dysbiosis.

We included fecal samples from 10 patients enrolled during regular out-patient visits. Control cohorts were selected from an array of previously published cohorts and matched to the CLL cohort on criteria including mean age, residency, year of sampling, and sampling methods. Using shotgun metagenomic sequencing and taxonomical profiling, we assessed fecal microbiome composition, diversity, and dynamics. Bioinformatic analyses were performed following the state-of-the-art methods as well as using innovative tailor-made approaches.

Ten patients diagnosed with CLL delivered 12 stool samples between January 2016 and October 2018. Sampling was prior to treatment for eight patients and after treatment for four patients, thus, two patients, were sampled both before and after treatment. No patients received antibiotics within four months prior to sample collection. For an overview of patient and control baseline characteristics, and patient treatment see Supplemental Figure S1A-C. Changes in relative abundance of 10 major bacterial genera in patients sampled both before and after treatment are demonstrated in Figure S1D.

We observed high intra-variability among the CLL microbiomes (Figure 1A). At the genus level, Bacteroides was the most abundant genus in six out of the 12 CLL samples. Additionally, there was a trend of Bacteroides acquiring bacterial dominance (>30% relative abundance) in 5 out of 12 samples. Several other bacterial genera such as Parabacteroides, Prevotella, and Acinetobacter also acquired bacterial dominance among CLL patients. We next assessed microbiome diversity in CLL patients and healthy controls. Richness (the observed number of species) was lowest for patients with CLL and was significantly different compared to each of the two control cohorts (Observed: CLL vs C1: median, 53 vs 69; P = .00057; CLL vs C2: median, 53 vs 73; P = 6.8e−05; Figure 1B). Patients with CLL also showed lower alpha diversity compared to the control groups when assessed by Shannon index (CLL vs C1: median, 1.90 vs 2.90; P = 2.1e−05; CLL vs C2: median, 1.90 vs 2.75; P = .00057) and InvSimpson index (CLL vs C1: median, 4.18 vs 11.94; P = 4.3e−05; CLL vs C2: median, 4.18 vs 10.05; P = .00057;
The two control cohorts also demonstrated a significant difference in diversity between one another (C1 vs. C2: Shannon: median, 2.90 vs. 2.75; P = .00057; InvSimpson: median, 11.94 vs. 10.05; P = .017). In addition to reduced diversity, a difference in the specific microbial composition between the CLL cohort and the two control cohorts was observed: At the phylum level, we focused on the differences in Bacteroidetes, Firmicutes, Proteobacteria and Actinobacteria as they comprised 95% of the total bacterial content in CLL patients, on average. The distribution of these 4 phyla across each cohort highlights a significantly higher abundance of Bacteroidetes and Proteobacteria relative to the controls (Figure 1C). In contrast, both control groups showed greater proportions of Firmicutes and Actinobacteria. 

At the family level, Bacteroidaceae, Prevotellaceae, Clostridiaceae, Lachnospiraceae, and Ruminococcaceae were the most abundant five families among both CLL patients and controls (Figure 2A). Bacteroidaceae were present in high proportions in the CLL cohort. Interestingly, a major difference was also observed between the samples from patients with CLL and the healthy controls with lower relative abundance of Lachnospiraceae and Ruminococcaceae taxa, while the dissimilarities between the control groups were small. We further aimed to identify groups of bacterial taxa differentially abundant between patients with CLL and controls. Analyzing the composition of microbiomes based on the relative abundance data at the genus level, we detected Bacteroides, Sutterella, and Parabacteroides, to be overrepresented in CLL relative to the average microbiome (abundance of a taxon across all CLL and healthy samples). In contrast, we identified a group of taxa including Bifidobacterium, Anaerostipes, and nine other bacterial genera to be underrepresented among patients with CLL as compared to controls (Figure 2B). Bearing in mind the complexity of bacterial communities, we also sought to characterize groups of bacteria co-occurring across all CLL and healthy samples. We calculated proportional co-occurrence coefficient for all pairs of bacterial genera across the CLL and healthy cohorts. When we visualized the relationships between 23 most strongly proportional taxa, three clusters were formed (Supplemental Figure S2A). Abundance/depletion of the proportional bacterial taxa across all samples revealed clear discrimination between CLL patients and healthy controls (Supplemental Figure S2B, Supplemental Table S1). An overlap of the two analytical methodologies indicated lack of bacteria mainly from Lachnospiraceae and Ruminococcaceae families among the CLL samples. In particular, lower abundances of Anaerostipes, Bifidobacterium, Blautia, Coprococcus, Dorea, Eubacterium, Ruminococcus, and Streptococcus were observed in CLL samples when compared to controls. The two approaches also indicated higher abundances of Bacteroides and Parabacteroides in CLL compared to controls. 

While canonical metrics have revealed statistically significant differences between CLL and healthy cohorts, we note that controls are also significantly different from one another across several metrics. Thus, we set out to answer whether there exists a healthy-diseased axis such that healthy patients from both control cohorts co-vary and are distinct from CLL patients. To do this we characterized the structure of covariance in each cohort by their centroids in PCA-space (Figure 3A). We assessed the similarity between cohorts by calculating the angles between their centroids within this reduced multidimensional space (interpretation of angles in Figure 3C). On average, CLL gut profiles were inversely proportional, i.e. dissimilar, to C1 & C2 cohorts (∠CLL,C1 = 140.1°± 38.5°, purple; ∠CLL,C2 = 168.9° ± 11°, green; Figure 3B). We also observed an acute relationship, i.e. similarity, between healthy cohorts (∠C1,C2 = 51°± 43.9°). These results highlight the presence of a strong inverse axis between diseased and healthy states that is defined by the angles between centroids of the CLL and the two control groups.
The lower fecal diversity observed in the CLL cohort is in line with previous findings of reduced bacterial diversity in other inflammatory conditions including IBD, type 1 and 2 diabetes, and obesity,\textsuperscript{10,11} as well as hematologic malignancies.\textsuperscript{6,12} We suggest that the decreased diversity/increase in Bacteroidetes and Proteobacteria observed in the CLL cohort could imply a general loss of bacteria, but may also indicate a loss of complexity for the remaining microbiome.

Most of the bacterial genera in the microbial signature depleted in patients with CLL belonged to Lachnospiraceae and Ruminococcaceae family. Members of the Lachnospiraceae and Ruminococcaceae families are among the main producers of short chain fatty acids (SCFA), which are known to modulate the surrounding microbial environment and to directly interact with the host immune system.\textsuperscript{13} Interestingly, Blautia (Lachnospiraceae family) and Ruminococcus (Ruminococcaceae family) were also identified with higher abundance among controls than among patients with CLL. High abundance of intestinal Blautia has been associated with improved survival upon graft-versus-host disease,\textsuperscript{14} and is together with Ruminococcus often underrepresented in feces samples from patients with colorectal cancer.\textsuperscript{15}

In conclusion, despite low sample size, the CLL microbiome demonstrated lower microbiome diversity and lower enrichment of SCFAs-producing bacterial taxa when compared to healthy controls. We hypothesize that the overabundance of bacteria from the Bacteroidetes phylum together with depletion of Lachnospiraceae and Ruminococcaceae family bacteria might play a role in – or is observed due to – immune dysfunction among CLL patients. This microbiome signature is warranting validation and refinement in larger CLL cohorts including patients needing treatment, patients assigned to watch and wait, patients with other hematological malignancies, as well as other healthy cohorts. This as a focus of ours in a follow up study will hopefully lay foundation for defining microbiome characteristics for hematological malignancies and microbiome signatures discriminating subgroup of patients with CLL.
References


Figure 1. Gut microbiome composition and diversity in CLL patients and controls. A) The relative abundance of bacterial genera in all 10 CLL patients (12 samples). Bacterial genera whose abundance was < 1.5% in a sample were grouped as 'Others'. Sequences that could not be assigned to a genus were grouped as 'Unclassified'. Taxa having zero counts across all samples were removed prior to all analyses. If the sample was taken after treatment, the treatment regimen is indicated by a corresponding shape on the top of each bar and described in the legend. Bacterial abundance was visualized using stacked bar plots from R package ggplot2. Unambiguously assigned genera: 1) [Rhodospirillum/Lactobacillus/Azospirillum]; 2) [Enterobacter/Escherichia/Klebsiella/Serratia]; 3) [Tidjanibacter/Alistipes]. B) Fecal diversity in CLL samples and healthy samples at genus level (alpha diversity measures: observed number of genera, Shannon and Inverse Simpson indexes). In box plots, box edges represent the 25th and 75th percentiles, the center line shows the median and whiskers extend from the box edges to the most extreme data point. Data beyond the end of the whiskers are plotted individually as dots. The p-values (adjusted for multiple testing with the Benjamini-Hochberg (BH)) obtained upon Wilcoxon rank-sum tests are indicated, values < 0.05 were considered significant. Ns, P > 0.05; *, P < 0.05; **, P < 0.01; ***, P < 0.001; ****, P < 0.0001. C) Relative abundance of four major bacterial phyla forming the microbiota in CLL cohort. Box plots are constructed as described in B.

Figure 2. Relative and differential abundance of bacterial families and genera. A) Relative abundance of five most abundant bacterial families in CLL and healthy feces samples. Box plots are constructed as described in 1B. Ns, P > 0.05; *, P < 0.05; **, P < 0.01; ***, P < 0.001; ****, P < 0.0001. B) Differential abundance of bacterial genera between CLL and the merged control cohorts. Bacterial genera are color coded according to its higher taxonomic rank - family. The y-axis W value represents number of times the null hypothesis (H0: the average abundance of a given taxon is equal across cohorts) was rejected for a given taxon. The x-axis value represents the centered log-ratio (clr)-transformed mean difference in abundance of a given taxon between the CLL and healthy groups with respect to average abundance of a given taxon. Positive value at the x-axis indicates bacterial genera being differentially abundant in controls, negative value indicates bacterial genera being differentially abundant in CLL cohort. Only bacterial genera with null hypothesis rejected in >70% of cases and clr mean difference -/+ 1 are labelled. The analysis and volcano plot visualization were done in R by implementation of Analysis of Compositions of Microbiomes (ANCOM).

Figure 3. Analysis of covariance by principal component analysis (PCA). To assess similarity between the CLL microbiome and the control microbiome in a multidimensional space, a PCA analysis was performed. A) The biplot illustrates the distance between the CLL cohort and the two control cohorts in terms of a 2-dimensional representative plot of 1000 iterations run on the original dataset (n=12 per cohort totaling 36 cases per cohort per iteration), delimited by principal component (PC) 1 and PC2. The large symbols (centroids) represent the mean PC score from each cohort. The PC score for each individual is plotted relative to their position along each of the PCs. The biplot shows vectors (black) pointing to the centroid of each cohort, as well as the individuals of each cohort (CLL – red triangles, C1_FR – purple squares, C2_AUS – green circles). Colored contour maps represent the density and distribution of individuals grouped by cohorts. The biplot is overlaid with a protractor-like plot displaying degrees from 0 to 180°. The angles between 2 cohorts were calculated as the angles between vectors pointing to centroids of individual cohorts ($cosθ = \frac{a \cdot b}{|a||b|}$), with CLL being always
positioned at 0°. Although C1_FR show certain overlap with CLL, note that the patients with CLL are distinctly clustered from both healthy control cohorts. B) The protractor-like plot represents all angles identified over the 1000 iterations between CLL-C1_FR and CLL-C2_AUS. The mean of all centroid vectors per cohort is drawn as a thick line with a white symbol at its end. The standard deviation (SD) is visualized by arrows of a color corresponding to the cohort, on the outside of the plot. C) The protractor-like plot provides interpretation of the angles on healthy-diseased axis. An angle between vectors is interpreted as an approximation of the correlation and the similarity between the cohorts’ variables; i.e., the C1_FR cohort has a dissimilar composition with weaker correlation with the CLL cohort, whereas the C2_AUS control cohort is inversely correlated to the CLL cohort.
Figure 3

A

B

C

Different composition and/or no correlation.

Dissimilar composition
Weak correlation

90°

Orthogonal axis

Similar composition
Proportional content

45°

135°

Similar composition
Inversely proportional

CLL @ 0°

Healthy-Diseased axis

180°

Same composition
Positive correlation

Same composition
Negative correlation
Supplemental Material

The Gut Microbiome in Patients with Chronic Lymphocytic Leukemia

Supplemental Figure S1. Patient and controls characteristic.

Supplemental Figure S2. Heatmap visualization of the highly proportional bacterial genera and microbial signature abundance.

Supplemental Table S1. Microbial signature of CLL identified by phi proportionality-clustering method.
**Supplemental Figure S1. Patient and controls characteristic.** A) Swimmer plot illustrating the clinical course of 10 CLL patients. The treatment regimens (black shapes), sampling time points (blue crosses) and survival status are shown in this swimmer plot. Patients 2 and 8 were sampled at two time points (Sample IDs: 2.1, 2.2 and 8.1, 8.2, respectively). In total, 8 samples were collected before treatment initiation out of which 2 were delivered at the day of treatment start; and 4 samples were collected after treatment initiation (highlighted by grey bars). Follow-up time differs for each patient and continues beyond the 130 days as indicated by arrow heads. B) CLL patient characteristics. All data shown here represent the time of diagnosis, except for Diagnosis-1st Treatment and Diagnosis-Death. Abbreviations: y, years; d, days; M, male; F, female; 1st, first-line; Diagnosis-1st Treatment, days from diagnosis to first-line treatment; Diagnosis-Death, days from diagnosis to death; NA, not available; SCLC, small cell lung cancer; BDM, bendamustine; IBR, ibrutinib; G, obinutuzumab; RTX, rituximab; VEN, venetoclax; Interm., Intermediate; B2M, beta-2 microglobulin; IGHV, immunoglobulin heavy chain variable region genes; U-CLL, unmutated IGHV-status; M-CLL, mutated IGHV-status; TP53, TP53 Gene - Tumor Protein P53; FISH, fluorescence in situ hybridization. C) CLL and healthy control groups characteristics. All data are no. (%) unless otherwise stated. IQR, interquartile range. D) Change in relative abundance (%) of the 10 most abundant genera in CLL samples. The samples are grouped based on collection timepoint (pre- or post-treatment). The arrows indicate increase (green) or decrease (red) in relative abundance of individual bacterial genera in the paired samples. Only change in relative abundance larger than 1% is marked. Color code of bacterial genera correspond to color code in Figure 1A. Samples collected from the same patient at 2 timepoints (paired-samples) are highlighted in light blue.
Supplemental Figure 2

A

Phi correlation coefficient

Clusters color code
- Cluster 1
- Cluster 2
- Cluster 3

B

Relative abundance (clt transformed)

CLL (N=12)  CONTROL 1 France (N=12)  CONTROL 2 Austria (N=12)

Relative abundance (clt transformed)
Supplemental Figure S2. Heatmap visualization of the highly proportional bacterial genera and microbial signature abundance. A) Hierarchical clustering of bacterial genera showing strong proportionality in terms of co-occurrence (phi (\(\phi\)) proportionality coefficient < 0.4) to any other bacterial genera from CLL and healthy samples. Clustered heatmap was built using hierarchical clustering with the \(\phi\) correlation coefficients as distance metrics. Tree cutting at second level resulted in the 3-cluster structure. The \(\phi\) statistic\(^1\) represented by \(\phi\) correlation coefficient indicates strength of proportionality between two variables (the closer the \(\phi\) to zero, the stronger the proportionality between two observed taxa). B) Heatmap of relative abundance of bacterial taxa from A) in CLL and healthy control groups. Clustering performed in A) was reused to cluster the bacterial genera here. Data used for visualization were clr-transformed. Upper: Twelve samples were randomly selected from each control group for better visualization. Lower: All samples from healthy controls were used for visualization.

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### Supplemental Table S1. Microbial signature of CLL identified by phi proportionality-clustering method

Table showing microbial signature divided into groups ‘CLL’ and ‘DEPLETED IN CLL’ based on three-cluster structure from Supplemental Figure S2 and the fold change. Group: Microbial signature group based on fold change in relative abundance between CLL and both control groups. Subgroup: Clusters of bacterial taxa corresponding to clusters in Supplemental Figure S2. Bacterial genera/family: Taxonomical classification. Identified by ANCOM: ✓ indicate that individual bacterial taxa were identified as differentially abundant by ANCOM (Figure 2B) independently from the phi (φ) proportionality analysis. Median relative abundance: Median relative abundance of individual bacterial taxa in individual cohorts, values are shown in percentages (%). Fold change: Fold change of individual bacterial taxa between CLL and control cohorts; calculated from median relative abundance. Taxa identified by an overlap of the two approaches (ANCOM and phi (φ) proportionality analysis) are considered as a potential microbiome pattern in the CLL patients.

<table>
<thead>
<tr>
<th>Group</th>
<th>Subgroup</th>
<th>Bacterial genera</th>
<th>Bacterial family</th>
<th>Identified by ANCOM (Fig. 2B)</th>
<th>Median relative abundance (%)</th>
<th>Fold change</th>
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<td></td>
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<td>CLL</td>
<td>CONTROL 1 France</td>
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