Evaluation of an effective detection and quantification method for particular microorganisms by comparing NGS-based metagenome profiling data

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Abstract

Metagenome profiling research using next-generation sequencing (NGS), a technique widely used to analyze the diversity and composition of microorganisms living in the human body, especially the gastrointestinal tract, has been activated, and there is a growing interest in the quantitative and diagnostic technology for specific microorganisms. According to recent trends, quantitative real-time PCR (qRT-PCR) are still of considerable technique in detecting and quantifying bacteria associated with the human mouth, nasal cavity, and pharynx due to analytical cost and time burden of NGS technology. Here, based on NGS metagenome profiling data produced by utilizing 100 gut microbiota samples, we conducted a comparative analysis of identifying for five bacterial genera proportions (Akkermansia, Bacteroides, Bifidobacterium, Phascolarctobacterium, and Roseburia) within same metagenomic DNA samples through qRT-PCR assay in parallel. Genus-specific primer, targeting the particular gene of each genus for qRT-PCR assay, allowed a statistically consistent quantification pattern with the metagenome profiling data. Furthermore, results of bacterial identification through Sanger validation demonstrated the high genus-specificity of each primer set. Therefore, our study suggests that an approach to quantifying specific microorganisms by applying qRT-PCR method can compensate for the concerns (potential issues) of NGS while also providing efficient benefits to various microbial industries.

Key Points

1) The quantitative real-time PCR (qRT-PCR) is an suitable detection and quantification tool that can counterpart the 16s metagenomics profiling technique.

2) The genus-specific primer designed by investigating a target site (or gene) for a microorganism of interest through large-scale data comparative analysis helps solve the mosaic problem of multiple conserved and variable regions of 16s rRNA

3) For bacterial identification through qRT-PCR, nucleotide sequence confirmation via Sanger sequencing is important for designing a genus-specific primer and developing a quantification method.

Introduction

The high-throughput nucleic acid sequencing technology (Next-Generation Sequencing; NGS) has developed rapidly over the past 15 years, applied to various molecular genetics studies (Gu et al. 2019; Slatko et al. 2018; Stenson et al. 2017). This technology is not only read whole-genome sequences for various species but also widely applied in the molecular diagnostic field, such as prediction of particular human disease through identifying biomarker genes (Okamura et al. 2019; Rim et al. 2017). In addition, advances in NGS technology have recently brought new changes to microbiological research (Cao et al. 2017; Deurenberg et al. 2017; Mallia et al. 2017). In past microbiological studies, microbial identification was possible through the conventional Sanger sequencing using bacterial 16S rRNA gene or fungal ITS (Internal Transcribed Spacer) regions only for culture-dependent microorganisms (De Filippis et al. 2018). However, the NGS technology, which can read multiple microbial genomes (be defined as ‘metagenome’) in specimens, allowed the classification of unculturable microorganisms also (Yarza et al. 2014). This approach to classifying microbiome through metagenome sequencing technology (e.g., whole-metagenome sequencing and 16S rRNA sequencing) has been used to understand complex microbial communities and biological interaction (Feng et al. 2018; Hemmat-Jou et al. 2018). Among the several metagenome sequencing technologies, reading the 16S rRNA gene is generally used as a phylogenetic marker to classify bacterial taxonomy (Theis et al. 2019).

The 16S rRNA gene (approximately 1500 bp) consists of the nine hyper-variable regions (V1-V9) interspersed among the high-conserved sequences. Because the nine variable regions are a powerful tool that can distinguish the diversity between bacterial strains, the universal primer pairs, including a single or a combination of these regions (e.g., V1V2, V3V4, V4, and V5V6 regions on the 16S rRNA gene), is used for 16S rRNA sequencing (Schierfer et al. 2018; Sperling et al. 2017; Walters et al. 2016). However, there is an issue that reading partial 16S variable regions is difficult to classify between some bacterial species with high-sequence similarity (Gao et al. 2017; Jeong et al. 2021). To overcome this bacterial misclassification problem, some companies (PacBio, Oxford Nanopore, and Loopsq) involved in NGS-based sequencing technology are developing novel metagenome analysis platforms that read the full-length 16S rRNA gene to classify microorganisms (Fujioishi et al. 2020; Karst et al. 2021; Metras et al. 2020). These developments of continuous metagenome research have led to efforts to establish an association between microbiota (e.g., inhabiting in intestinal mucosa or oral cavity) and human health (Mohajeri et al. 2018; Young 2017). Recently, human gut health has been checked by profiling the gut microbiome composition through feces samples (Abrahamson et al. 2017; Liang et al. 2020; Williams et al. 2019).

The human gastrointestinal tracts are composed of more than 100 trillion different microorganisms such as bacteria, archaea, and viruses, creating a complex microbial ecosystem (Rinninella et al. 2019). For the last ten years, large-scale gut microbiome studies such as the Human Microbiome Project (HMP) and MetaHIT explained that many human inflammatory diseases, obesity, and neurological diseases are associated with ‘dysbiosis’ (Gevers et al. 2012; Heiman and Greenway 2016; Nash et al. 2017). The dysbiosis, which is defined to the imbalance of normal microbiota inhabiting the gut, is mainly caused by bad eating habits, stress, and antibiotics. This phenomenon results in an increasing proportion of intestinal harmful bacteria and yeast associated with various human diseases (Baj et al. 2019; DeJong et al. 2020; Losso 2021). In this respect, NGS-based microbial classification has been applied to various microbiome studies, resulting in significant biomedical findings. Still, the metagenome sequencing technology has some issues such as a high-cost burden and time-consuming if the purpose is intended to detect just specific microorganisms (O’Sullivan et al. 2013). For these reasons, the real-time PCR (also denoted as quantitative real-time PCR—qRT-PCR) is usually used to confirm rapid microbial detection through target gene quantification from clinical samples taken from participants (Pereira et al. 2017; Wang et al. 2011). Indeed, many companies and researchers involved in microbial molecular diagnosis are attempting to overcome the issue of high-sequence similarity between bacterial species arising from the 16S rRNA gene by developing specific primer pairs for using qRT-PCR equipment.

In this study, we investigated the concordance between qRT-PCR and NGS technology to confirm the efficiency of microbial quantification for genus level associated with the human health. Since NGS is currently not suitable as a method for identifying species-level microorganisms, we have confirmed the
similarity of the two technologies at the genus level. Based on NGS metagenome profiling data produced by utilizing 100 gut microbiota samples, we conducted a comparative analysis of quantification accuracy for five bacterial genera (Akkermansia, Bacteroides, Bifidobacterium, Phascolarctobacterium, and Roseburia) through qRT-PCR assay in parallel. Genus-specific primer targeting the particular gene (such as transcription termination/anti-termination protein; nusG) with one of the genera allows relatively similar quantification results with the 16S V3-V4 metagenome profiling data. Additionally, we cross-validated the genus-specificity of the designed target primers via Sanger sequencing with qRT-PCR products as template DNA.

**Materials And Methods**

1) **Human stool samples collection**

One hundred human stool samples were collected using the OMNigene-GUT stool swab kit (DNAGenotek, Canada) from healthy adult men and women participating in the Korean Gut Microbiome Database Project of the Korea Food Research Institute (KFRI, South Korea). All stool samples were then stored at -80°C for the experimental downstream processing. The sampling was carried out with the prior informed consent of all participants before this study began. None of participants affected the research results, such as taking medication before the study began. The clinical sample collection in this study (stool samples collection) was approved by the ethics committee of Theragen Bio (Theragen Bio, South Korea) Institutional Review Board (IRB Protocol Number: 700062-20180905-JR-005-01). All clinical experiments applied to this study were carried out according to the guidelines and regulations of the declaration of Helsinki.

2) **Metagenomic DNA extraction**

Total metagenomic DNA (mDNA) from 100 stool samples was isolated using the QIAamp DNA Stool Mini Kit (Qiagen, Germany), and all experimental processes were performed in accordance with the optimal protocols provided on the DNA extraction kit. The quality check of all isolated mDNA was conducted using a Bioanalyzer (Agilent 2100, USA) at the Center for Bio-medical Engineering Core Facility (Dankook University, South Korea). All mDNA samples were then stored at 4°C until the following process.

3) **Illumina 16S V3-V4 amplicon sequencing library preparation and sequencing**

A total of 100 metagenome sequencing libraries were prepared according to the Illumina 16S amplicon sequencing library construction workflow (Illumina, USA). The Illumina platform targeted an area containing the V3-V4 hyper-variable region of the bacterial 16S rRNA gene. The PCR amplification of the target region was started immediately after the mDNA was extracted. The 16S V3-V4 amplicon was amplified using KAPA HiFi Hot Start Ready Mix (2×) (Roche, Germany). For this purpose, a pair of V3-V4 target-specific universal primers recommended by Illumina were used. The primer sequences were as follows: 16S 341F forward primer is 5'-TCGTCGGCAGCGTCAGATGTGTATAAGAGACAGCCTACGGGNGGCWGCAG-3' and 16S 806R reverse primer is 5'-GTCTCGTGGGCTCGGAGATGTGTATAAGAGACAGGTCTTAGGAGATGTGATAAGACAGGAGACCAGCTACGGGNGGCWGCAG-3'. After the PCR amplification, the clean-up process of all PCR products was conducted using the AMPure XP beads (Beckman Coulter, USA). And then, additional PCR amplification was performed to add multiplexing indices and Illumina sequencing adapters using the Nextera XT Index Kit (Illumina, USA). The final PCR products were then purified once again using the AMPure XP beads. After the amplicon library construction, the 16S metagenome sequencing was carried out using the paired-end 2 × 300 bp Illumina MiSeq protocol (Illumina MiSeq, USA; (Wu et al. 2016)).

4) **Bacterial genus-specific primer design methods**

Genus-specific primers for qRT-PCR amplification were designed to identify and absolute quantify the particular five bacterial genera from mDNA. The overall process for primer design in this study is as follows.

1. **Bacterial genera and target gene selection.** Based on 16S V3-V4 metagenome sequencing profiling data, five human intestinal bacteria at the genus level associated with health or disease were listed. Particular genes to detect each bacterial genus through the qRT-PCR were then selected.

2. **Sorting of NCBI annotation information.** The National Center for Biotechnology Information Database (NCBI DB) was used to obtain sequence information about the selected genes to target each bacterial genus. First, the 'Identical Protein Groups' category was used to extract a comprehensive summary table that explains the gene annotation information on the NCBI DB (e.g., nucleotide and protein accession number, organism at species and strain level, etc.). Next, the ambiguously classified information on the table, such as hypothetical protein, was excluded from the list. Then, repetitive information about the randomly selected bacterial species (included strain level) belonging to each genus was filtered. Finally, all 'Protein ID' on the filtered list were isolated to obtain the amino acid sequence information needed to convert to gene sequence.

3. **Extraction of coding sequence information.** The Batch Entrez open web bioinformatics tool linked with NCBI DB (www.ncbi.nlm.nih.gov/sites/batchentrez) was used to convert amino acid information about specific genes into coding sequences (CDS) in the form of the FASTA format.

4. **Multiple sequence alignment and selection of target-specific regions.** To find the consistent sequence regions for qRT-PCR primer design, the multiple sequence alignment of the CDS information was performed using the BioEdit 7.2v software.

5. **In silico test.** The experimental suitability (Tm value, GC%, and potential for primer-dimer to form, etc.) for qRT-PCR of the selected primer pairs were checked using the Oligo calc and Oligo Analysis open web tool (http://biotools.nubic.northwestern.edu/; http://www.operon.com/tools/oligo-analysis-tool.aspx). Next, the NCBI Nucleotide BLAST (https://blast.ncbi.nlm.nih.gov) was used to confirm the primer specificity for the targeted bacterial genus.

5) **Bacterial quantification using qRT-PCR**
The qRT-PCR was conducted to quantify the five bacterial genera frequencies from the isolated 100 mDNA samples. First of all, double-stranded DNA concentration within all mDNA samples was measured using the Qbit Fluorometer 4.0 v at the Center for Bio-medical Engineering Core Facility (Dankook University, South Korea) and 1X dsDNA HS Assay kit (ThermoFisher Scientific, USA). Then, all template mDNA was then normalized to an identical concentration (10 ng/ul) via dilution by using distilled water. Next, a standard curve for 100 diluted DNA samples was calculated to confirm that the template DNA concentration used in qRT-PCR was consistently normalized. In this step, the cycle threshold (Ct) value for amplicon quantity in qRT-PCR measured using the bacterial 16S V4 primer pair (515F and 806R) was applied. These Ct values were reflected from the 10-fold serial dilution (10^-1, 10^-2, and 10^-3 dilution) samples for each formerly normalized mDNA sample. Finally, the bacterial genera frequency within 100 template mDNA was confirmed using the StepOnePlus™ Real-Time PCR (ThermoFisher Scientific, USA) at the Center for Bio-medical Engineering Core Facility (Dankook University, South Korea) and Quantispeed SYBR No-Rox kit (PhileKorea, South Korea). Each qRT-PCR primer annealing temperature condition applied at this step is as follows; *Akkermansia*: 65 °C, *Bacteroides*: 65 °C, *Bifidobacterium*: 62.5 °C, *Phascolarctobacterium*: 62 °C, and *Roseburia*: 65 °C. Each summary table (Supplemental Table S6), which included the Ct value of five bacterial genera, was used to compare NGS-based bacterial frequency data.

6) Sanger sequencing

The Sanger sequencing was conducted to verify the molecular specificity and experimental accuracy of the five primer pairs designed for this study. First of all, the ten qRT-PCR amplicon samples (Top five and bottom five of the measured Ct value) were sorted from each qRT-PCR summary table (Supplemental Table S6) for the Sanger validation. The selected amplicon samples were then purified using the FavorPrep™ GEL/PCR Purification Kit (Favorgen, Tiwan). Next, molecular cloning through ligation and transformation process were carried out using TOPcloner™ TA Kit (Enzymics, South Korea) and DH5α chemically competent *E. coli* (Enzymics, South Korea) to obtain template DNA containing amplicon sequence. A colony PCR amplification using the M13 primer pair (M13F: 5’-GTAAAACGACGGCCAG-3’; M13R: 5’-CAGGAAACAGCTATGAC-3’) was conducted to check whether the target sequence was inserted in the plasmid cloning vector. The colony PCR products were once again purified through the FavorPrep™ GEL/PCR Purification Kit (Favorgen, Tiwan). The Sanger sequencing with purified products as template DNA was carried out using the ABI 3500XL Genetic Analyzer sequencer (ThermoFisher Scientific, USA) at the Center for Bio-medical Engineering Core Facility (Dankook University, South Korea). The ABI and FASTA format data generated after sequencing were used as input data in the NCBI Nucleotide BLAST tool for bacterial genus identification.

7) 16S V3-V4 data processing and microbial community analysis

The 16S V3-V4 sequencing reads were demultiplexed using the split_libraries_fastq.py function in QIIME2 metagenome analysis pipeline and sequences were quality trimmed using the Divisive Amplicon Denoising Algorithm 2 (DADA2) pipeline in R (version 3.3.2) with the parameters; EE=2, TruncL=c (200, 180) and q=10. The set of unique 16S V3-V4 DNA sequences, referred to as amplicon sequence variants (ASVs), were then inferred using DADA2 and an ASV table of quality trimmed using the Divisive Amplicon Denoising Algorithm 2 (DADA2) pipeline in R (version 3.3.2) with the parameters; EE=2, TruncL=c (200, 180) and q=10. The set of unique 16S V3-V4 DNA sequences, referred to as amplicon sequence variants (ASVs), were then inferred using DADA2 and an ASV table of read counts per ASV per sample was generated. ASV taxonomy were taxonomically classified using the sklearn-based Naive Bayes classifier with the SILVA 138v 16S rRNA database.

8) Statistical analysis

An association analysis of the bacterial genera proportions data within 100 samples, measured from the qRT-PCR assay and 16S metagenome sequencing results, was conducted through a statistical test using the Spearman correlation test (Fig. 4; Table 3). The Spearman correlation's statistical significance evaluated the statistical similarity between the two methods was denoted as asterisk (*) if P-value < 0.05 and R value (Spearman's value) > 0.5.

Results

1) Selection of five bacterial genera from 16S metagenome analysis data

This study confirmed the consistency of the relative proportion pattern within the samples of five particular bacterial genera between the 16S metagenome profiling data and qRT-PCR quantitative results (Fig. 1). We collected human stool specimens from 100 healthy Korean (adult men and women). Then, we extracted mDNA from the 100 specimens and simultaneously checked the DNA quality (e.g., DNA degradation, concentration, and purity) for accurate microbial quantification. To select bacterial genus for quantifying frequency in the specimens, we successfully prepared the Illumina 16S V3-V4 short-read amplicon sequencing libraries using the 100 mDNA samples, and NGS-based 16S metagenome sequencing was performed. As a result of 16S metagenome sequencing, the average demultiplexed reads count generated from 100 samples was 133,288, of which 65,892 were filtered through the DADA2 pipeline with non-chimeric reads. (Supplemental Table S1). After 16S V3-V4 sequencing data processing, the total number of bacterial ASV taxonomy classified from the SILVA 138v 16S rRNA gene reference database was 6,902 (with more 70% classification confidence threshold), of which 338 were classified at the genus level (Supplemental Table S2). Based on the ASV taxonomy classification results at the genus level, we selected the *Akkermansia*, *Bacteroides*, *Bifidobacterium*, *Phascolarctobacterium*, and *Roseburia*. These five genera are the composition of human gut microbiota and are probiotic bacterial strains closely related to obesity, dysbiosis, and Inflammatory Bowel Disease prevention. (Gomez-Gallego et al. 2016; John and Mullin 2016; Liu et al. 2018; Russell et al. 2011; Wang et al. 2020). They are also beneficial bacteria that contribute significantly to the production of the short-chain fatty acid (SCFA; e.g., acetic, butyric, and propionic acid), which is key to human health control, such as increasing immunity, maintaining intestinal homeostasis, and preventing fat accumulation in the body (Markowiak-Kopec and Silzewska 2020). Considering these interesting bacterial features, we applied these five selected genera to compare the relative proportions of particular bacterial load within the samples, measured using the qRT-PCR assay and NGS frequency data.

2) Bacterial genus-specific primer design

We selected the five bacterial taxa (*Akkermansia*, *Bacteroides*, *Bifidobacterium*, *Phascolarctobacterium*, and *Roseburia*) at the genus level based on the formerly performed 16S metagenome profiling data. To quantify the bacterial load within the samples using the qRT-PCR assay, we designed genus-specific
primer pairs to detect these five genera (Table 1). Considered that the high-sequence similarity of the 16S rRNA gene internal hyper-variable region would be an obstacle to detecting the specific genus, we used sequence information on certain bacterial gene domains annotated on the NCBI reference database (Supplemental Table S3; show detailed in the Materials and Methods section). First of all, we applied the sequence information of *nusG* gene, which encodes a transcription termination/antitermination protein essential for bacterial transcription (Lawson et al. 2018). For the *Bifidobacterium* and *Akknemansia*, it was challenging to find consistent regions in the multiple sequence alignment results between the bacterial species belonging to the genus. Therefore, we reselected the transaldolase and D-alanine D-alanine ligase (*ddl*) gene sequences as a target-specific gene for *Bifidobacterium* and *Akknemansia*. In the case of *Bifidobacterium*, we referred to previous studies that *Bifidobacterium* species has a gene coding region capable of expressing at least 14 types of transaldolase that can be differentiated using the protein electrophoresis method (Requena et al. 2002). Additionally, we selected the *ddl* gene as a target gene of *Akknemansia*, considering that it is an essential factor for bacterial cell wall synthesis (Ellsworth et al. 1996). Finally, we conducted an *in silico* test using the NCBI Nucleotide BLAST web tool to validate the bacterial genus-specificity of five primer pairs designed for this study (Supplemental Fig. S1). Although some *in silico* test results showed classification results for microorganisms other than the targeted bacterial genus (e.g., *Raphanus sativus*, *Rodentibacter pneumoniae*, and *Acetobacterium woodii*), we determined that it would not affect our qRT-PCR assay because they were eukaryotic or not human gut bacteria (Banhani 2017; Fingas et al. 2019; Bache and Pfenning 1981).

3) Quantification and Normalization of metagenomic DNA

Before estimating a particular bacterial load within the human fecal samples, we normalized the bacterial DNA concentration, which was included in the 100 mDNA samples, through the standard curve calculation using the qRT-PCR (Fig. 2; Table 2; Supplemental Table S4). We used 10-fold serial diluted mDNA samples (10⁻¹, 10⁻², and 10⁻³ dilution from 10 ng/ul concentration of dsDNA per each sample) and bacterial 16S V4 universal primer (515F and 806R) for this qRT-PCR assay. Theoretically, considering that the 16S rRNA gene is the most conserved region on the bacterial genome, we have determined that it is appropriate to apply the 16S V4 specific-primer pair to the internal bacterial genomic DNA quantification of the mDNA samples (Frank et al. 2008; Tran et al. 2017). As a result of the qRT-PCR assay, we confirmed that the average Ct values about all 10⁻¹, 10⁻², and 10⁻³ diluted samples for the V4 amplification were measured at 27.31, 23.26, and 19.85, respectively. In addition, we could also confirm that the standard deviation (SD) and the coefficient of variation (CV) value of each Ct value for the three dilution factors were averaged 0.55 and 2.41 percent. Calculating the standard curve based on these measurements, the R² value of the trend line connecting Ct values for all 10-fold diluted samples was about 0.97. Additionally, we found that the R² value of the trend line was close to 1 when calculating the standard curve based on the average Ct value. These results showed that our systematic quantification challenge for the particular genera was conducted with the bacterial DNA samples measured and normalized by comprehensive calculation methods.

4) Parallel comparison of qRT-PCR and 16S metagenome profiling data

We compared the relative bacterial proportion similarity of the five genera within the human gut microbial community between the qRT-PCR assay using formerly normalized 100 mDNA samples (10 ng/ul per sample) and the NGS-based 16S metagenome profiling data (Fig. 3; Fig. 4; Table 3; Supplemental Table S5; Supplemental Table S6). In order to this comparative approach, we calculated the potential bacterial frequency (1/2^{Ct value} and proportion of each genus within the 100 samples using the Ct value exported by qRT-PCR assay. The multiple comparison data in the Fig. 3 showed that relative distribution of the five bacterial genera proportions within each sample was visualized by the bacterial abundance bar plot and cumulative curve graph. Comparing the qRT-PCR assay with NGS proportions data in parallel, we found that the relative proportion patterns of the five bacterial genera were remarkably consistent between the two different quantification methods. We highlight that four bacterial (*Akknemansia, Bifidobacterium, Phascolarctobacterium, and Roseburia*) genus present in low proportions in 100 samples of human gut microbiota also confirmed significant relative proportions of the quantification results. Additionally, we verified an association between bacterial proportions data measured from two different quantification methods using the Spearman correlation statistical analysis to supplement these parallel comparison results. As the statistical analysis result, we could confirm that the Spearman correlation values for all bacterial genus groups were significantly calculated (P-value < 0.05; Spearman's rho value > 0.5). Considering that the Spearman analysis evaluates statistically positive similarity if the correction value (Spearman's rho value) is 0.4 or higher, we could confirm the significant similarity of the relative bacterial proportion measured between the NGS and qRT-PCR assay (Ramsey 1989). Thus, our comparative results showed that the five genus-specific primer pairs have a high-binding sensitivity and specificity for the target mDNA.

5) Verification of primer specificity

We performed Sanger sequencing of the qRT-PCR amplicon products to confirm the of the bacterial genus-specificity of five primer pairs designed for this study, and verified the sequencing results by BLAST (Basic Local Alignment Search Tool) (Fig. 5; Table 4; Supplemental Table S7). First, we conducted a ligation process to insert qRT-PCR amplicon products with top 5 and bottom 5 Ct values measured per each bacterial genus-group as a TOPO TA cloning plasmid vector for transformation. We applied the TOPO TA vector with M13 primer regions (M13F and M13R; approximately 200 bp) because the input library length for high-quality Sanger sequencing results is at least 400 bp (Kchouk et al. 2017). After the transformation process, we isolated five white colonies (potentially transformed component cells) per each LB culture medium and performed direct colony PCR using the M13 primers. Next, we performed the Sanger sequencing to confirm the sequence information of the qRT-PCR amplicon. When the Sanger sequencing results were confirmed through the BLAST search using the blastn parameter, we found that the taxonomy definition rate with the NCBI DB was almost 100% in the four groups (high and low proportion groups of the *Akknemansia, BacteroidesPhascolarctobacterium*, and *Roseburia*). For the *Bifidobacterium*, the bacterial taxonomy definition rates of high and low groups were 92%, and 60%, respectively. Although some BLAST results of the *Bifidobacterium_Low* groups were resulted to be undefined with NCBI DB, we could find that the relative proportion for 100 people calculated from the NGS frequency and qRT-PCR Ct value was almost similar. As a result of Sanger validation, the qRT-PCR and NGS proportion data shown in Supplemental Table S6 of the *Bifidobacterium_Low* group samples for which the NCBI nucleotide BLAST search was not valid (less than 60% of bacterial taxonomy definition rate per each plate) were as follows; Bif_Mi_01 (Candidate 50): 0.01 and 1.80, Bif_Mi_02 (Candidate 77): 0.01 and 0.00, Bif_Mi_03 (Candidate 83): 0.01 and 0.00. In this regard, this issue found at the *Bifidobacterium_Low* group was likely to result in low transformation efficiency in the TA cloning process due to i) the too low bacterial frequency within the samples and ii) high G+C content
(approximately 63 mol%) in the transaldolase gene coding region of the *Bifidobacterium* genome, rather than misbinding of the specific primer pairs that occurred during the qRT-PCR amplification (Freitas and Hill 2018; Park et al. 2019; Serafini et al. 2012). Considering that identification of the target bacterial genus was possible in all Sanger validation results of the selected genus groups reflecting the high and low genus proportions within each sample, we confirmed that the five specific primer pairs designed for this study are sufficient to be applied to bacterial detection and quantification using the qRT-PCR assay.

**Discussion**

Since the Human Genome Project, high-throughput sequencing-based microbiome research projects such as the HMP and MetaHIT consortium have emphasized the importance of identifying the association between microorganisms and human diseases. In particular, these projects explained that the causes of various physical diseases are closely related to the "dysbiosis" phenomenon, in which normal intestinal microbiota is unbalanced (Consortium 2010; Gevers et al. 2012). Accordingly, various probiotic therapy-associated industries and researchers have classified the human gut microbial composition using NGS technology to identify the relative intestinal proportions of beneficial and harmful bacteria and their biological effects (Sharifi-Rad et al. 2020; Shukla 2017). However, NGS technology's high analytical cost burden and time consumption are considered important issues in microbiome-associated health care industries. Furthermore, the microbial misclassification problem at the species level caused by the high sequence similarity within the 16S rRNA gene is also regarded as a challenge to be addressed (Callahan et al. 2021, Jeong et al. 2021). Therefore, some researchers have suggested that running experiments in parallel with qRT-PCR, which enables accurate and rapid detection and quantification of the target genes, is a solution to compensate for these issues of the NGS technology. Because the high fluorescence sensitivity of the equipment enables accurate detection of specific microbial species in the sample, the qRT-PCR assay is still used as a 'gold standard method', especially in the molecular microbial diagnostics and probiotics fields (Masco et al. 2007; Shehata et al. 2019).

Through parallel comparison analysis, we evaluated that the qRT-PCR technique represents an excellent alternative to existing NGS-based 16S V3-V4 metagenome sequencing methods as it enables reliable detection and quantification (for the gut-associated microflora: *Akkermansia*, *Bacteroides*, *Bifidobacterium*, *Phascolarctobacterium*, and *Roseburia*). Since quantifying the gut microbiome composition of healthy subjects is necessary to establish a baseline against which microbiome changes can be detected in pathological conditions, we performed normalization to the initial microbial density in the V3-V4 hyper-variable region of the 16S rRNA gene. We minimize errors in specific microbial populations through this process by quantifying the total load of internal standards and 16S rRNA genes in the meta-samples (Tran et al. 2017). The process of absolute quantification of internal standards as well as in bacterial quantitation systems utilizing qRT-PCR will avoid sacrificing high rates due to abundance and amplification bias in amplicon sequencing that can occur in NGS-based metagenome sequencing and is essential to maintain reproducibility (Zemb et al. 2020). Although the 16S rRNA gene primer is specifically designed and optimized for qRT-PCR or NGS, it is recommended to consider the potential bias and the miss annealing due to sequence similarity between specific bacterial strains for qRT-PCR. In our study, the qRT-PCR primers target CDS sites of a gene (e.g., bacterial housekeeping genes such as *nusG* or *ddl*) representing a particular microbial genus to ensure high sensitivity and reduce biased quantitative results by sequence similarity. In addition, to evaluate the accuracy and sensitivity of the designed genus-specific primer, we selected five amplicon samples with the high and low frequency generated after qRT-PCR amplification, respectively, and confirmed Sanger sequencing validation. As a result, almost all samples where genus-specifically amplified amplicons were generated, suggesting how specific our target primers are and how respectable they are for particular genus quantification. We assumed that several undefined samples through Sanger sequencing validation might not be identified due to the reduced PCR cloning efficiency with the high G+C contents of the target CDS or the minimal DNA content of the *Bifidobacterium*. Nonetheless, the evaluation of the specificity and sensitivity of primers used for the quantification of microorganisms using Sanger sequencing validation suggests that it is an essential task as a standard research procedure for clinical diagnostic research and industrial development. As a result of cross-checking the correlation of the measured bacterial proportions between two different quantification methods through Spearman correlation statistical analysis, it was also verified that the qRT-PCR method was sufficient and accurate for quantitative analysis of specific microbial genera in meta-samples. Therefore, we propose an efficient system in fields requiring rapid quantification of indicator microorganisms required in medicine, agriculture, marine biology, and etc.

In conclusion, we confirmed through a parallel comparative analysis with 16S metagenome sequencing data that the qRT-PCR method can complement some issues (such as high analytic cost burden or time consumption) of the NGS technology, which applies to various microbial industries. Our findings showed that the relative bacterial proportions pattern of the five genera within samples, measured via the qRT-PCR assay performed under normalized mDNA concentration conditions, was statistically similar to 16S metagenome sequencing data. Additionally, we cross-validated bacterial genus-specificity of the five primer pairs designed for the qRT-PCR assay through the Sanger sequencing and NCBI BLAST test for bacterial identification. In this respect, we affirmed that a qRT-PCR method is one of appropriate tools for identifying the relative proportions of particular microorganisms within the sample. Furthermore, we suggest that applying qRT-PCR to specific microbial validation in the NGS-based microbial diagnosis industry can compensate for the concerns of NGS technology by providing economical, fast, and accurate services to consumers in terms of turnaround time.

**Declarations**

**Author's contributions**

K.H. designed and supervised the project; C.C., S.M., Y.O. and J.J. performed experiments; K.Y., T.H., W.C., M.L., A.Y., S.M., and J.J. analyzed the data; Y.A., K.H., S.M., and J.J. wrote the manuscript. All authors read and approved the final manuscript.

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Data Availability

The datasets generated during the current study are included in this published article and its supplementary information file and are available from the corresponding author on reasonable request.

Ethical approval

All clinical samples (stool samples) were collected from healthy adult men and women participating in the Korean Gut Microbiome Database Project of the Korea Food Research Institute (KFRI, South Korea). The human experiment in this study (samples collection) was approved by the ethics committee of Theragen Bio (Theragen Bio, South Korea) Institutional Review Board (IRB Protocol Number: 700062-20180905-JR-005-01)

Conflicts of interest

The authors have no conflicts of interest to declare that are relevant to the content of this article.

Consent to participate

Not applicable.

Consent to publication

Not applicable.

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Tables

Table 1. Overall information of bacterial genus-specific primer set

| Bacterial taxon         | Rank | Target gene | Foward primer (5'-3')                  | Reverse primer (5'-3')       | Tm °C (F/R) | GC % (F/R) |
|-------------------------|------|-------------|---------------------------------------|------------------------------|-------------|------------|
| Akkermansia             | Genus| ddl         | CTTCGTGCTGAAATCAACACC                 | CGATAATTCGGCTATTITTCGC       | 62.1/59.2   | 50/3!      |
| Bacteroides             | Genus| nusG        | GGTGCCCTCTGACAATTACAG                 | CAATGATACACTGAATCCGCT        | 60.5/60.1   | 55/4!      |
| Bifidobacterium         | Genus| Transaldolase| AAGGGCATCTGGCTGTAACCG                | GAGACGAGAAGAGGACGA           | 59.5/60.5   | 58/5!      |
| Phascolarctobacterium   | Genus| nusG        | TCCCTGGTATGTTGCTTGTAGAG              | CAGTCAAAGGAATCCTTTAGTA       | 60.9/59.2   | 43/3!      |
| Roseburia               | Genus| nusG        | AAATACCCGTGGTGTTACCG                 | GTGTCTCCCTCTGTAAAGTCA        | 58.4/59.5   | 50/4!      |

Table 2. Average standard curve calculation results using qRT-PCR assay

| Dilution factor          | Average *Ct value | *SD value | *CV value | Target gene       |
|--------------------------|-------------------|-----------|-----------|-------------------|
| 10^-3 from 10 ng         | 27.31             | 0.49      | 1.78      | 16S rRNA V4 region|
| 10^-2 from 10 ng         | 23.26             | 0.52      | 2.24      | 16S rRNA V4 region|
| 10^-1 from 10 ng         | 19.85             | 0.64      | 3.21      | 16S rRNA V4 region|

*Ct value: Cycle threshold value; *SD value: Standard deviation value; *CV value: Coefficient of variation value

Table 3. Statistical result of the Spearman correlation test between two different quantification methods

| Bacterial genus         | *R value | Spearman P-value | Spearman's Sig. |
|-------------------------|----------|------------------|-----------------|
| Akkermansia             | 0.89562266 | 2.98E-36         | ***             |
| Bacteroides             | 0.62412241 | 0                | ***             |
| Bifidobacterium         | 0.8538906  | 1.51E-29         | ***             |
| Phascolarctobacterium   | 0.6444568  | 4.67E-13         | ***             |
| Roseburia               | 0.51864254 | 3.25E-08         | ***             |

*Sig.: Statistical significance; *R: Spearman’s rho value
Table 4. Bacterial identification result by Sanger sequencing

| Bacterial genus  | Defined bacterial taxon counts in NCBI database | Defined bacterial taxon rates (%) of Sanger validation |
|------------------|-----------------------------------------------|------------------------------------------------------|
|                  | Major Top 5 (Ct value) | Minor Top 5 (Ct value) | Total | Major Top 5 (Ct value) | Minor Top 5 (Ct value) | Total |
| Akkermansia      | 25                     | 25                     | 50    | 100.00                | 100.00                | 100.00 |
| Bacteroides      | 25                     | 24                     | 49    | 100.00                | 96.00                 | 98.00  |
| Bifidobacterium  | 23                     | 15                     | 38    | 92.00                 | 60.00                 | 76.00  |
| Phascolarctobacterium | 25                 | 25                     | 50    | 100.00                | 100.00                | 100.00 |
| Roseburiae       | 25                     | 25                     | 50    | 100.00                | 100.00                | 100.00 |

Figures

Figure 1

Experimental introduction in this study and schematic workflow of genus-specific primer design method for qRT-PCR assay. Using stool samples obtained from 100 healthy adults, we performed a comparative analysis to evaluate the accuracy of microbial quantification by two different molecular technologies. (a) The overall experimental workflow of the qRT-PCR assay and NGS-based 16S V3-V4 metagenome sequencing. (b) Schematic diagram showing the process to design the genus-specific primer set to quantify the proportions of the five selected bacterial genera within each sample using qRT-PCR.
Figure 2

Standard curve calculation to confirm normalization of the mDNA concentration used for qRT-PCR analysis. Each double-stranded mDNA sample diluted with a three step 10-fold serial dilution (10-1, 10-2, and 10-3) was standardized at 10 ng/ul for standard curve calculation. The graph's x-axis indicates the concentration of each 10-fold diluted mDNA used in standard curve calculation, and the y-axis shows the Ct value measured from the qRT-PCR. (a) Standard curve graph representing the Ct value of every samples with 10-fold serial diluted mDNA concentration (R2 = 0.97). (b) Standard curve graph representing the average Ct value of each concentration of the diluted mDNA sample (R2 = 1.00).

Figure 3

Parallel comparison of the interest five bacterial genera proportions measured from two different quantification methods. Multiple overlaid connected line graphs show the quantitative identity of each bacterial genus measured by both methods. The graph's x-axis indicates the 100 samples (denote as 'candidate'), and the y-axis indicates the relative proportion value for each particular bacterial genus. The bar plot shown on the left side represents an average abundance of five bacterial genera calculated from 16S metagenomic profiling analysis.

Spearman correlation test between qRT-PCR and NGS proportion

Correlation Coefficient Shows Strength & Direction of Correlation

Strong Weak Weak Strong

-1.0 -0.5 0.0 +0.5 +1.0

Negative Correlation Zero point Positive Correlation

Taxon (Spearman's Rho value)

Akkermansia (P<0.05) Bacteroides (P<0.05) Phascolarctobacterium (P<0.05) Roseburia (P<0.05)
Figure 4

Spearman correlation scatter plot showing the relationship of each bacterial relative proportion value measured by qRT-PCR assay and 16S V3-V4 metagenome sequencing data; *** = Spearman P-value < 0.001, R = Spearman's rho value.

Figure 5

Verification of the specificity and accuracy of primers for quantifying the five bacteria genera using Sanger sequencing. (a) The schematic diagram shows the overall experimental process for TA cloning of each genus-specific amplicon, bacterial gene transformation, Sanger sequencing validation, and sequence confirmation through NCBI nucleotide BLAST search. (b) The confirmation of the nucleotide sequence of the genus-specific amplicons, showing five high and low frequencies, respectively, is shown in the mosaic plot. The sky-blue and red colors denote the BLAST test identified the appropriate target bacterial genus name and not working, respectively.

Supplementary Files

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