Oral Metagenomic Analysis Techniques

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The modern era of microbial genome analysis began in earnest in the 2000s with the generalization of metagenomics and gene sequencing techniques. Studying complex microbial community such as oral cavity and colon by a pure culture is considerably ineffective in terms of cost and time. Therefore, various techniques for genomic analysis have been developed to overcome the limitation of the culture method and to explore microbial communities existing in the natural environment at the gene level. Among these, DNA fingerprinting analysis and microarray chip have been used extensively; however, the most recent method of analysis is metagenomics. The study summarized examined the overview of metagenomics analysis techniques, as well as domestic and foreign studies on disease genomics and cluster analysis related to oral metagenome. The composition of oral bacteria also varies across different individuals, and it would become possible to analyze what change occurs in the human body depending on the activity of bacteria living in the oral cavity and what causality it has with diseases. Identification, isolation, metabolism, and presence of functional genes of microorganisms are being identified for correlation analysis based on oral microbial genome sequencing. For precise diagnosis and treatment of diseases based on microbiome, greater effort is needed for finding not only the causative microorganisms, but also indicators at gene level. Up to now, oral microbial studies have mostly involved metagenomics, but if metatranscriptomic, metaproteomic, and metabolomic approaches can be taken together for assessment of microbial genes and proteins that are expressed under specific conditions, then doing so can be more helpful for gaining comprehensive understanding.

Key Words: Dental caries, Metagenomics, Microbiome, Oral

Introduction

The microbiome is defined as “all microorganisms and genes in a given environment”1). It is possible to study the role of whole microbial communities efficiently through recent advances in next-generation sequencing (NGS). Approximately <1% of microorganisms are known to be culturable under general laboratory conditions2,3). Microbes living in the human body are also unculturable, and it is estimated that approximately 20~60% of microbes cannot be cultured although it depends on the body parts4). As most microbes are not culturable (the great plate count anomaly), methods for analyzing microbial flora through culturing are no longer used5,6).

Most human microbiomes are composed of bacteria; therefore, techniques focusing on bacterial community analysis have been developed and reported in several studies. It is now possible to identify almost all species because the database for bacteria is well-established with 16S rRNA gene as the phylogenetic barcode, which also requires a taxonomy database that is accurately organized7).

The rapid development of genetic information and data analysis technologies such as NGS developed in the 2000s has led to active metagenomic studies that analyze approximately 3.9×10¹³ symbiotic microbial communities at microbial gene level rather than through culturing.

In terms of the diagnosis and improvement of the health and treatment of disease, the human body and symbiotic microorganisms are being studied together. Research into the microbiome enables us to identify the principles of human health and disease from a new perspective and devise more fundamental and effective new treatments.

Therefore, the present study aims to investigate the analytical methods used in metagenomic studies and the
Effects that characterization of oral metagenomics could have on periodontal disease and dental caries.

Materials and Methods

This study is a review of research articles on the microbial techniques for genomic analysis and information on oral microorganisms and genomes. In stage 1, we designed and planned research content related to the characteristics of microbial techniques for genomic analysis and information on oral microorganisms, and in stage 2 we performed a literature search. In stage 3, based on the selected literature, we collated the research contents. In stage 4, we objectively interpreted and derived conclusions from the analyzed data. The period for the literature search and organization of results was from 1st January to 30th April, 2019. For the literature search, we used PubMed, KISS (Koreanstudies Information Service System), DBpia, and RISS (Research Information Sharing Service), with the search terms, ‘microbiome’, ‘ngs’, ‘oral’, ‘metagenomic’, ‘subgingiva’, ‘caries’, and ‘saliva’.

Results

1. Technology and analysis of microbiome

1) Metagenomics

In metagenomics, DNA is extracted directly from the crude source without culturing microbes or employing a microbial isolation technique. The extracted metagenome contains a mixture of genomic DNA of all the microbes present in the sample. Its community structure is analyzed at the gene level through NGS. The development of metagenomics is interlocked with the commercialization
of NGS, which began in 2007\(^9\). Previously, large-scale metagenomics-based studies were difficult to standardize because DNA sequencing was expensive. The purpose of metagenomics is to first determine and analyze the microbes present in the sample and second, to identify the metabolic processes and functional genes present in the sample. The shotgun (whole-genome shotgun) metagenomics approach is mainly used to view the functional gene. There is an advantage in viewing the entire metagenome, including the composition of functional genes; however, there are disadvantages too, such as the high cost of sequencing and difficulty in sequence identification\(^{10}\). Conversely, to determine individual microbial species present in the diverse microbial community, an amplicon metagenomics method is used that selectively amplifies the marker gene for species identification and then analyzes the sequence of the amplification product. Thus, the effort and high-cost for analysis can be reduced greatly (Fig. 1)\(^11,12\).

2) Next-generation sequencing

NGS is a DNA sequencing method that is an improvement over the existing Sanger method, and it can acquire a large number of sequences in a short time at a relatively low cost. The first NGS was achieved through Roche’s 454 pyrosequencing. The method used at that time divided the entire gene into sequence fragments of 200 bp each, which was read, and then arranged in a single line to reconstruct the whole genome code from the original 3 billion codes\(^13\). Typical NGS technologies include Roche’s 454 pyrosequencing; Illumina’s sequence by synthesis; Pacific BioSciences’ single molecule, real-time sequencing; and Oxford Nanopore’s nanopore sequencing technologies\(^14,15\). These technologies are far superior in throughput and cost than the existing Sanger method and are superior in read length and accuracy. NGS technologies allow the analysis of all fields of genomes across DNA, RNA, and epigenetics\(^16\). This kind of genome analysis provides a new paradigm of technology to all fields of biology and medicine by enabling the diagnosis and target treatment together (Table 1)\(^17\).

3) 16S rRNA amplicon sequencing

The 16S rRNA is a ribosomal RNA that constitutes a ribosome that synthesizes proteins. It has a length of 1,542 nucleotides and has a conserved region common to all species and a hypervariable region capable of classifying a specific species. The 16S rRNA gene is amplified, and the sequence is analyzed using a massive parallel method; it is a sort of an acting barcode that can identify various kinds of microbes because each microbe has a different sequence of hypervariable region. There is little mutation among the same species, but some mutations are found among different species. These differences are compared to identify the closeness between the species. If more than 97% of the 16S rRNA sequences are identical, they are the same species, if more than 94% of them are identical, they are the same genus, if more than 90% of them are identical, they are the same family, and if 85% of them are identical, they are classified as the same order\(^5,18\).

The 16S rRNA gene is used as a marker gene to analyze the bacterial communities, and the primer used depends on the type of sequencing device. The 27F and 518R primers are used to amplify the mutation site V1 ~ V3 portions when using GS Junior (Roche). The 454 technique reads a sequence backwards from the reverse primer, resulting in a sequence of 400 ~ 450 bp in length, starting at V3 and reaching the edge of V1. In contrast, if MiSeq v3 (Illumina) is used for sequencing, 318F and 806R primers are used to amplify the V3 ~ V4 portions. A sequence of at least 400 bp in length is produced by combining a pair of sequence 300 bp in length, from both forward and reverse sides, into one through paired-end sequencing\(^19\).

Eren et al.\(^20\) used 16S rRNA gene sequence data from 9 areas in the oral cavity, and 493 oligomer types were identified from V1 ~ V3 data, and 360 oligomer types were identified from V3 ~ V5 data. These types of oligomers were associated with taxonomic names at the species level by comparison with the Human Oral Microbiome Database (HOMD). They have discovered oligomer types that were sometimes differentiated by a single nucleotide that has different distributions between the oral sites and the individuals\(^20\).
### Table 1. Comparison of Next-Generation Sequencing Platforms

| Machine (manufacturer) | Chemistry | Run time | Gb per run | Current, approximate cost (US$) | Advantages | Disadvantages |
|------------------------|-----------|----------|------------|-------------------------------|------------|---------------|
| HiSeq 2000/2500 (Illumina) | Reversible terminator | 23 h | 0.7 | 500,000 | Cost-effectiveness | High error rate in homopolymers, Long run time, Minimal hands-on time, Very short read lengths, Instrument not available at time of writing (available end 2012) |
| MiSeq (Illumina) | Proton detection (Illumina) | 27 h | 1.5 | 125,000 | Cost-effectiveness, Short run times, Appropriate throughput for microbial applications, Minimal hands-on time | High error rate in homopolymers, read lengths too short for efficient assembly, Read lengths too short for efficient assembly |
| HiSeq 2000 (Illumina) | Proton detection (Illumina) | 23 h | 0.7 | 750,000 | Cost-effectiveness, Short run times, Appropriate throughput for microbial applications, Minimal hands-on time | Long error rate in homopolymers, read lengths too short for efficient assembly, Read lengths too short for efficient assembly |
| Ion Proton (Life Technologies) | Proton detection (Life Technologies) | 2 h | 145,000 | 125,000 for compulsory server | Appropriate throughput for microbial applications, Minimal hands-on time | High error rate in homopolymers, read lengths too short for efficient assembly, Read lengths too short for efficient assembly |
| PacBio RS (Pacific Biosciences) | Reversible terminator | 20 min | 3 per d | 750,000 | Simple sample preparation, Low reagent costs, Very long read lengths | High error rate, Expensive installation, Difficult installation |
| Ion Personal Genome Machine (Life Technologies) | Proton detection (Life Technologies) | Up to 200 | 1.5 | 80,000 (including OneTouch and server) | Appropriate throughput for microbial applications | High error rate in homopolymers, read lengths too short for efficient assembly, Read lengths too short for efficient assembly |
| Ion Proton (Life Technologies) | Proton detection (Life Technologies) | Up to 10 (Proton I chip) or up to 100 (Proton II chip) | Up to 10 (Proton I chip) or up to 100 (Proton II chip) | 145,000+75,000 (Proton I chip) or 125,000 (Proton II chip) | Appropriate throughput for microbial applications, Minimal hands-on time | High error rate in homopolymers, read lengths too short for efficient assembly, Read lengths too short for efficient assembly |
| Ion Personal Genome Machine (Life Technologies) | Proton detection (Life Technologies) | Up to 200 | 1.5 | 80,000 (including OneTouch and server) | Appropriate throughput for microbial applications | High error rate in homopolymers, read lengths too short for efficient assembly, Read lengths too short for efficient assembly |
| Ion Proton (Life Technologies) | Proton detection (Life Technologies) | Up to 10 (Proton I chip) or up to 100 (Proton II chip) | Up to 10 (Proton I chip) or up to 100 (Proton II chip) | 145,000+75,000 (Proton I chip) or 125,000 (Proton II chip) | Appropriate throughput for microbial applications, Minimal hands-on time | High error rate in homopolymers, read lengths too short for efficient assembly, Read lengths too short for efficient assembly |

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Table 1 shows a comparison of next-generation sequencing platforms, including their machine manufacturers, chemistry, read length, run time, Gb per run, current approximate cost, advantages, and disadvantages. The table includes platforms such as HiSeq 2000/2500 (Illumina), MiSeq (Illumina), HiSeq 2000 (Illumina), Ion Proton (Life Technologies), PacBio RS (Pacific Biosciences), and Ion Personal Genome Machine (Life Technologies). Each platform is compared in terms of its chemistry (Reversible terminator, Proton detection), run time (23 h, 27 h), Gb per run (0.7, 1.5), current approximate cost (500,000, 125,000), advantages (Cost-effectiveness, Short run times), and disadvantages (High error rate in homopolymers, Long run time, Minimal hands-on time).
4) Shotgun metagenomics sequencing

Shotgun metagenomics sequencing is a method of analyzing functional genes in microbial communities. This is an innovative technology that mosaics the DNA extracted from the sample by a super computer after cutting it into arbitrary fragments and analyzing the sequences. The original data generated by the sequencing device contains errors that occur during the experiment and the sequencing process. The cause of errors, such as innate or inherent errors of polymerase during polymerase chain reaction (PCR), production of chimeric sequences between heterologous DNA, and homopolymer errors occurring during sequencing, are very diverse; the types of errors also depend on the experimental method or the sequencing device. These erroneous sequencing reads are filtered out, and then taxonomic assignments and diversity calculations are performed for the remaining reads. Each sequencing piece is assembled with contigs by joining the overlapped parts, and then the scaffolds are completed by listing the contigs in the order. This approach allows for the entire metagenome analysis, including the composition of complex functional genes, and has the advantage that all microbial organisms can be found at once. Currently, various bioinformatics analysis platforms have been developed, and among them, QIIME21), MOTHUR22), RPD23), and PlutoF24) are the most frequently used programs. These platforms perform separation, identification, and diversity calculations based on code, by filtering only the sequences suitable for analysis from the original data25).

2. Metagenomic community analysis of oral cavity

All techniques used for analyzing oral microbes have strengths and limitations. Based on the first selectively collected human microbial community data, the HOMD, information on microorganisms and genomes was obtained26). Subsequently, the 16S rRNA database of the core oral microorganisms was created, which is a key phylogenetically selected core database27).

The HOMD includes 619 taxa in 13 phyla, as follows: Actinobacteria, Bacteroidetes, Chlamydiae, Chloroflexi, Euryarchaeota, Firmicutes, Fusobacteria, Proteobacteria, Spirochaetes, SR1, Synergistetes, Tenericutes, and TM7. Although the oral microbial phyla have been expanded to 15, only 6 phyla account for 95% or more of the sequence, as follows: Actinobacteria, Bacteroidetes, Fusobacteria, Proteobacteria, and Spirochaetes28).

Jorth et al.29) confirmed the changes in the composition and gene expression of oral microbiota in healthy subjects and patients with periodontitis through expression patterns of the transcriptome. In a study conducted using samples collected from nine healthy subjects and nine patients with periodontitis, 160,000 genes were compared. The differences in the composition and gene expression of microbiota between the healthy subjects and patients with periodontitis were revealed through massively parallel RNA-sequencing, and a difference in metabolism between the two groups was observed. Conversely, the expression of genes involved in the metabolism of individual species within each microbiome was greatly diverse for each individual. In addition, disease-related microbiomes showed conservative changes in metabolism and toxic gene expression29).

Frias-Lopez and Duran-Pinedo30) used metatranscriptome analysis for periodontal plaque in laboratory experiments and revealed that there was a rapid change in the gene expression pattern of microbiota of healthy individuals upon the addition of periodontal pathogens to the dental plaque of multi-bacterial species. Duran-Pinedo et al.31) used subgingival microbiomes of six healthy subjects and seven patients with periodontal diseases and analyzed the in situ genome-wide transcriptome. The metabolic activities that occurred during the disease were found to be iron acquisition, lipopolysaccharide synthesis, and flagella synthesis.

Wang et al.32) described the key microbes associated with periodontitis through metagenomic sequencing. From the sequencing data of a total of 10 healthy individuals, 10 variables shared by 11 bacterial species were identified33). Based on several literature, major genera in the mouth include: Streptococcus, Veillonella, Granulicatella, Gemella, Actinomyces, Corynebacterium, Rothia, Fusobacterium, Porphyromonas, Prevotella, Capnocytophaga, Neisseria, Haemophilus, Treponema, Lactobacterium, Eikenella, Leptotrichia, Peptostreptococcus, Staphylococcus, Eubacteria, and Propionibacterium34).

In an integrated study conducted as part of the Human Microbiome Project (HMP) since 2012, samples were
collected from 10 different sites in the digestive tract of more than 200 healthy individuals to compare and analyze the composition of microbiomes. In this study, salivary microbiota of healthy oral cavity was similar on the throat, tonsil, and the back of the tongue. The dominant phyla identified were Firmicutes, Bacteroidetes, Proteobacteria, and Fusobacteria, and they accounted for 40%, 25%, 20%, and 10%, respectively. The dominant genera were found to be Streptococcus and Veillonella, and they accounted for 20% and 15% of the salivary microbiota, respectively. NGS has currently been used to examine the saliva of two adults in five data sets extracted from HMP, and more than 175 bacteria have been identified in oral saliva. Considering that the salivary microbiota is far more complex than expected, the prevalent oral microbiota of saliva is predicted to consist of approximately 900 different bacterial species.

In another study in 2011, saliva samples from 74 children were analyzed by pyrosequencing. The occupancy rate of Firmicutes decreased as the salivary microbiota decreased from deciduous teeth to permanent teeth, but the ratio of Bacteroidetes and Proteobacteria increased as well as the total bacterial diversity. Recent studies have reported that the salivary microbiota may be altered in patients with inflammatory bowel disease, emphasizing the dynamic association between microbiomes at different anatomical locations.

3. Metagenomic study in the field of dental disease

1) Periodontal disease

The virulence of the total microbiomes is increased by the core pathogen, Porphyromonas gingivalis, which interacts with secondary pathogens, like Streptococcus mitis (strain B6), and generally causes dysbiosis of the beneficial microbiota and inflammatory diseases. Every person has a group of individual-specific microorganisms that are essential for maintaining health and controlling disease. Griffen et al. found differences in the microbiomes between the healthy condition and periodontitis in all phylogenetic and specific microbiomes by using 16S multiple region pyrosequencing. Spirochaetes, Synergistetes, and Bacteroidetes were the dominant phyla in diseases, and a higher level of Proteobacteria appeared in healthy condition. Through this study, we were able to determine the association between P. gingivalis, Treponema denticola, and Tannerella forsythia and the disease. Abusleme et al. performed 454 pyrosequencing using 16S rRNA gene library and confirmed that Spirochaetes, Synergistetes, Firmicutes, and Chloroflexi were higher in the microbe of periodontitis, while Actinobacteria was higher in healthy condition. Kirst et al. investigated the subgingival microbiota of 25 patients with chronic periodontitis and 25 healthy subjects by using rRNA sequencing and compared them with the HMP. This study suggested that microbiota significantly changed with decreasing heterogeneity in periodontal disease. Two microbial communities were confirmed; one was Fusobacterium and Porphyromonas, which were associated with periodontitis, and the other one was Rothia and Streptococcus, which were associated with healthy condition.

Several studies have confirmed the presence of periodontal pathogens in saliva. A large-scale study in Finland used a 16S rRNA-based PCR method along with species-specific primers, where at least one of the periodontal pathogens, including T. forsythia, T. denticola, P. gingivalis, Campylobacter rectus, Aggregatibacter actinomycetemcomitans, and Prevotella intermedia, was found in 88% of samples in both groups of patients with periodontitis and healthy subjects. In another study using the 16S rRNA-based PCR method, six periodontal pathogens, including Prevotella nigrescens, A. actinomycetemcomitans, P. gingivalis, T. forsythia, P. intermedia, and T. denticola, was identified in the saliva samples of 41 children with healthy oral condition. P. nigrescens accounted for 80%, T. denticola for 32%, A. actinomycetemcomitans for 24%, and P. gingivalis for 12% of the pathogens. In a study in 2014, saliva samples of a total of 586 Danish adult patients, including 139 patients with periodontitis, and 447 healthy individuals were analyzed using the human oral microbe identification microarray (HOMIM) technique, and 12 phylotypes related to periodontitis were identified in saliva. A recent study using 16S rRNA pyrosequencing compared...
gingival plaque samples and saliva samples from 19 patients with periodontitis before and after periodontal therapy. This study showed that gingival microbiota was significantly different from salivary microbiota; the gingival microbiota significantly changed after periodontal treatment, but salivary microbiota was less affected by periodontal treatment53).

2) Dental caries

According to the report on microbes related to caries through pyrosequencing of dental caries in samples, *Streptococcus mutans* was not the dominant microbe in dental caries, but a complex microbiome was involved54). This supported the previous 16S rRNA sequencing study55,56) and the notion that dental caries were multiple bacterial diseases. In addition, through pyrosequencing, we found that oral bacteria are specific to the progression of caries57). *Streptococcus, Granulicatella*, and *Actinomyces* genera were significantly increased in children with severe dental caries58). The NGS technology, coupled with metagenomics, showed overexpression of functional genes acting as antimicrobial peptides or quorum sensing in individuals who did not experience dental caries, and thus, did not carry *S. mutans*54).

The analysis of the saliva of children aged 3～6 years and samples of gingival plaque using high-throughput barcoded pyrosequencing and PCR denaturing gradient gel electrophoresis according to the presence of dental caries resulted in the identification of 156 genera belonging to 10 phyla of the saliva. In addition, salivary microbiota was reported to be different from the gingival plaque microbiota, but the genera associated with patients with dental caries have not been reported59). Analysis of the saliva samples of 30 caries-active and 20 healthy Chinese children aged 6～8 years by using the HOMIM technique confirmed a total of 94 taxa in 30 genera belonging to the representative 6 phyla. We were also able to confirm that 8 and 6 taxa showed a significant difference in the saliva samples from children with dental caries and healthy children, respectively60). In an integrated study using a 16S rRNA gene amplicon or whole genome-based sequencing technique to analyze saliva samples from 19 patients with dental caries and 26 healthy subjects aged 18～22 years, salivary microbiota greatly varied in patients with dental caries compared with the healthy subjects61). In a recent study on the expression of functional genes, functional gene microarray; HuMiChip 1.0 was used to analyze saliva samples from 10 dental caries patients and 10 healthy persons aged 18～23 years, and we were able to confirm the association between the expression of functional gene related to disease and saliva samples from dental caries patients62).

**Conclusion**

Although studies on microbial genomes have been actively conducted worldwide, the progress has been very slow and many studies still focus on how microbial imbalances lead to pathologies in human body and on their diagnosis. Recent trends in molecular biochemistry have entered the field of omics, which is a comprehensive study of all biochemical molecules that target individual genes in organisms.

As studies on large genome-wide data have currently progressed into a new phase, studies that focused on whole genome analysis, such as transcriptome sequencing and epigenome sequencing, were conducted concurrently in various large-scale groups. However, there are still limitations, such as nonrandom sampling, disease complexity, absence of large biological database, and lack of breakthrough bioinformatics analysis methods.

With the rapid development of molecular diagnostic techniques, it is important to recognize that very common oral diseases, such as periodontitis and dental caries, are multifactorial diseases and also to integrate genomics and transcriptomics to identify individuals vulnerable to diseases63). In addition, it is necessary to focus on the functional aspect of microbiota by studying the microbiome and the complexity of the oral microbiota based on metagenomics.

**Notes**

**Conflict of interest**

No potential conflict of interest relevant to this article was reported.
Ethical approval
It is a reviewed paper so no deliberation required.

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