Oral and Human Microbiome Research

Sung-Kyun Chung†
Department of Dental Hygiene, Shinhan University, Uijeongbu 11644, Korea

In the past gut microbiome has been the main focus of microbiome research. Studies about the microbiome inside oral cavities and other organs are underway. Studies about the relationship between noninfectious diseases and periodontal diseases, and the negative effects of harmful oral microbes on systemic health have been published in the recent past. A lot of attention is being paid towards fostering a healthy oral microbial ecosystem. This study aimed to understand the roles and effects of the microbiome inside the human body can potentially help cure various diseases including inflammatory bowel diseases with no known cure such as Crohn’s disease, atopic dermatitis, obesity, cancer, diabetes, brain diseases and oral diseases. The present study examined technological trends in the correlation between the human microbiome and diseases in the human body, interactions between the human body’s immunity, the metabolic system, and the microbiome, and research trends in other countries. While it has been proven that human microbiome is closely correlated with human diseases, most studies are still in the early stage of trying to compare the composition of microbiomes between health and patient groups. Since the oral environment is a dynamic environment that changes due to not only food intake but also other external factors such as lifestyle, hygiene, and drug intake, it is necessary to continue in-depth research on the microbiome composition characteristics to understand the complex functions of oral microorganisms. Analyzing the oral microbiome using computational technology may aid in disease diagnosis and prevention.

Key Words: Gastrointestinal microbiome, Metagenome, Microbiome, Oral

Introduction

The Human Genome Project was regarded as a breakthrough technology in the 1990s, and people expected that it would provide a solution for a considerable portion of intractable diseases by analyzing all the base pairs in humans1). However, analysis of the database revealed that the human genome was much simpler than expected, and that this data alone is insufficient in explaining the complex biological processes in the human body. Following this, many scientists realized the necessity of studying symbiotic microbes in humans and their genes, and started a second genome project. They made an inventory of microbial genes and genomes in an effort to understand over 1,000 species of human microbiota, their 2~4 million genes, and the interaction between humans and these microbes2). The vast majority (95%) of the microbiota symbiotic with humans resides in digestive organs including the colon, while a large number of microbes also live in respiratory organs, reproductive organs, oral cavity, and on the skin3). As the microbiome is becoming the central focus in next-generation bio-healthcare, attempts are being made to exploit the microbiome for prevention of diseases and enhancement of human health, rather than simply identifying correlations and causations through experiments. This review summarizes the present knowledge regarding the microbiome and outlines recent microbiome research.

Introduction to the Microbiome

The microbiome is an ecosystem with humans, which consists of bacteria, viruses, and eukaryotic microbes such as fungi. There are variations between individuals in the
species that comprise the microbiota. However, the core microbes that play an indispensable role in keeping us healthy, exhibit considerable consistency, despite differences in individual characteristics and environmental conditions. Microbiome research began to gather momentum in the late 2000s when it was called ‘next generation genomics’, and findings from these studies establish that the microbiome has profound effects on not only human physiology, but also on human health and disease. For instance, the microbiome plays a role in monosaccharide absorption in the small intestine and fat buildup in the liver⁴. In particular, the human microbiome differs between individuals and experiences dynamic changes, depending on factors such as family, race, sex, age, diet, region, and lifestyle. Microbes inhabit our skin, oral cavity, teeth, respiratory organs, and reproductive organs, extract nutrients and energy from the food we eat, and help to repel or suppress pathogens⁶.

Microbiome is a compound made up of the words ‘microbiota’ and ‘genome’. It was first defined in 2001 as “the community of commensal (with no helpful exchange or pathogenesis), mutualistic and pathogenic microorganisms that share our body space”⁷. The 2006 study lead by Dr. Jeffrey Gordon from Washington University, that contends that lean and obese people have different gut microbiota, initiated widespread recognition of the microbiome as an essential element in human health⁸. In another study by Dr. Jeffrey Gordon, the feces of obese and lean mice were inserted into germ-free mice. Follow-up observations revealed that even with the same feed consumption, mice that had the feces of obese mice inserted into them gained an additional 50% of body fat in 2 weeks compared to those that received lean mice’s feces. They reported that the gut microbiota not only assists in digestion and nutrient absorption but also affects diseases like obesity⁹. Based on these study results, the US launched the Human Microbiome Project (HMP), anticipating that analyses of the human microbiota would provide clues regarding the mysteries surrounding human diseases¹⁰. The International Human Microbiome Project (IHMP) was formed in order to analyze and share the genetic information of specific microbes that cause diseases in citizens of each nation, and the project has been engaged in active research since then. Reports from various activities and research show that the microbiome is closely involved in nutrient absorption, regulation of drug metabolism, development of the immune system and brain, and infectious disease prevention¹¹.

The Role of the Human Microbiome

The human fetus develops in a microbe-free environment, but comes in contact with microbes as it passes through the birth canal during delivery, swallows maternal bacteria during breastfeeding, and via skin contact. Once the microbes settle, they develop into a unique microbiome for the individual human depending on a myriad of factors including the external environment and diet. The human microbiome enables us to digest and absorb nutrients we cannot digest alone, plays an active role in metabolism, and affects our immune system. Thus, its implications are not limited to intractable diseases like cancer or Crohn’s diseases, but are directly linked to general human health.

1. Nutrient absorption

Not all enzymes needed to digest the food we eat are produced in the human body. Around 4,000 species of microbes in the gut complement the function of enzymes by aiding in nutrient absorption, producing vitamins, and preventing inflammation. One of the roles of microbes is carbohydrate digestion. Gut microbiota break down many parts of proteins, lipids, carbohydrates, and other nutrients from the food before our body can absorb them. The microbial (produced during the digestion of consumed food) and human metabolites interact with other microbes, and play a role in diverse areas including the regulation of the immune system, metabolism, nervous systems, and brain and behavior development. The gut microbiota shows remarkable variations between individuals according to the type of food they consume and their dietary changes. Despite controlling for nutrient uptake, we found an association between gut microbiome composition and nutrient absorption profiles of individuals. Previously differences in nutrient uptake profiles were only attributed to genetic differences¹².
2. Immune system regulation

The concept of the human immune system, a system protecting our body from external pathogens, is expanding to include its interaction with and regulation of the human microbiome\(^{13}\). Microbiome function and population are regulated by immune cells and antibody activity. The microbiome is important in the development of lymphatic systems, which include the thymus and spleen\(^{14}\). Gut microbiome digestion of carbohydrates produces short-chain fatty acids (SCFA), increases production of interleukin (IL)-10, and reduces production of IL-6 and tumor necrosis factor-\(\alpha\). SCFA is a critical fatty acid which affects immune system regulation, vitamin production, and protects the intestinal mucosa\(^6\). Lack of gut microbiome settlement during the neonatal period hampers SCFA formation and leaves the infant vulnerable to autoimmune diseases, atopic dermatitis, asthma, and allergies\(^{14}\). Correlations between immune tolerance and the microbiome in patients with atopic dermatitis have been reported. The reduced diversity in skin Gamma-proteobacteria in patients with atopic dermatitis decreases IL-10 production, and the heightened proportion of Actinobacteria in the skin of healthy individuals induces an increased production of IL-10\(^{15}\).

3. Drug metabolism regulation

The microbiome plays a role in regulation of drug response, breaking down or modifying administered drugs or carcinogens in the metabolism of ingested substances. Through various metabolic and catalytic activities, it protects our body from drugs and carcinogens entering the body\(^{16,17}\). The microbiome influences various metabolic reactions including dehydroxylation, decarboxylation, dealkylation, and deamination, and impacts oxalate metabolism\(^{18-20}\). This indicates that metabolism of many drugs like anticancer and cardiovascular drugs, can be facilitated or inactivated by the microbiome, and different treatment regimens may be needed as the same drug may have different reactions among different patients due to their microbiome differences\(^{21}\). By harnessing the regulative capabilities of the microbiome in drug metabolism, we can provide customized prescriptions for patients with specific diseases based on a proper diagnosis of their genome and microbiome.

4. Prevention of infectious diseases

The microbiome functions as a barrier that blocks infection from external pathogens, and it is known that the microbiome balance can significantly alter our defensive capacity against pathogens. Microbiome imbalance weakens barrier function and causes mucosal damage in the intestinal tract, causing pathogens, toxins, and antigens in the intestine to enter the bloodstream and stimulate the immune system, thus leading to infections and autoimmune diseases. An excess of fast food and processed food, a Westernized diet, frequent alcohol consumption, and antibiotic use prevalent in urban lifestyles, disrupt gut microbiota diversity and cause gut microbiota dysbiosis, resulting in allergies, atopic dermatitis, and inflammatory diseases in the digestive system. In a study that compared travelers who contracted diarrhea during a trip with those who did not, researchers found that the presence of \textit{Lactobacillus ruminis} in higher proportions in a person’s microbiome was linked with lower diarrhea incidence and severity\(^{22}\).

**Microbiome and Diseases**

The human microbiome is an ecosystem consisting of various symbiotic microbes, and the discovery of its relationships with disease and health led to the possibility of using microbes in diagnosing and treating modern diseases without known origins.

1. Obesity and diabetes

Over 90% of bacteria living in the intestines of patients with obesity belong to the Firmicutes phylum, while Bacteroidetes constitute 3% of the population. Firmicutes and Bacteroidetes are gut microbes related to obesity; Firmicutes causes obesity while Bacteroidetes inhibits it\(^{8,23}\).

Inducing obesity in healthy mice through a high-fat diet changes its microbiome to an obese type, and transplanting the gut microbiome of obese mice into a germ-free mice facilitates fat accumulation in the mouse\(^{9,24,25}\).

Although causative relationships are unclear, there were more Clostridia in the microbiome of patients with type 2
diabetes than in healthy controls. Other studies found that butyrate-producing bacteria are reduced in patients with type 2 diabetes\textsuperscript{26}. The association between gut microbiota and diabetes is being investigated, and some studies reveal that the microbiota and its metabolites contribute to type 2 diabetes by affecting energy, carbohydrate metabolism, and inflammation\textsuperscript{27}).

2. Inflammatory bowel disease

Microbiome composition is a major factor that causes inflammatory bowel disease (IBD), the major types of which are ulcerative colitis and Crohn’s disease. Though the disease shows a wide range of symptoms and clinical courses, genetics and gut microbiome are thought to be common causes\textsuperscript{28}). Studies on the relationship between IBD and gut microbiome have found reduced gut microbiota diversity, an increase in Enterobacteriaceae, and a decrease in Firmicutes including \textit{Fecalibacterium prausnitzii}. When comparing patients with Crohn’s disease and ulcerative colitis with controls, patients with Crohn’s disease show decreased Firmicutes and increased Gammaproteobacteria, \textit{Escherichia coli} and \textit{Mycobacterium paratuberculosis}. An Enterobacteriaceae increase was characteristic of ulcerative colitis, and both Crohn’s disease and ulcerative colitis patients exhibited a decrease in \textit{F. prausnitzii}\textsuperscript{29}). Mutualistic gut microbes were found to be a play a role in increased immunological homeostasis and IBD suppression and treatment. Further research on the exact role of gut microbiota and immune response is expected to advance the development of diagnostic and therapeutic methods for IBD.

3. Cognitive impairments

Analysis into the gut microbiota of healthy mice and mice with Alzheimer’s disease confirmed that the two groups had different amounts and compositions of gut microbiota, and inserting the gut microbiota of mice with Alzheimer’s into a healthy rat caused a rise in beta-amyloid levels, which is considered to be the main causative factor of Alzheimer’s. Analyses to identify bacteria associated with Alzheimer’s revealed that there were less Firmicutes and Actinobacteria, and Bacteroidetes were more abundant. Based on a twin study report on dementia that linked tooth loss and dementia, the association between oral microbial composition and dementia was hypothesized\textsuperscript{30}). Two mechanisms explaining the association between oral microbiome and the brain, especially in Alzheimer’s disease, have been proposed\textsuperscript{31}). First, aging decreases microbiome-stabilizing oral saliva secretion, thus causing harmful microbes to overpopulate. Second, trigeminal and olfactory nerves may serve as a passage for oral microbes to invade the brain or transmit substances such as amyloids secreted by the microbes\textsuperscript{32}).

4. Cardiovascular disease

Arteriosclerosis is usually caused by plaque formation in the arteries. The plaques harden and eventually clog the arteries. However, some people with arteriosclerosis have no plaque, and toxic metabolites were detected inside their intestines. These toxic metabolites, such as trimethylamine produced by gut microbiota, are suspected to work in a manner similar to plaque\textsuperscript{33}). Cardiovascular risks including heart failure and stroke increase with increase trimethylamine levels. The gut microbiota generates trimethylamine from carnitine and lecithin, causing cardiovascular diseases\textsuperscript{33}). The oral microbiome is also known to affect cardiovascular disease. Together, these results suggest that everyday human microbiome management is important in order to prevent and alleviate a variety of diseases, including cardiovascular diseases.

Global Research Trends

The microbiome is receiving a lot of attention as a major global research interest in the age of the Fourth Industrial Revolution, following innovations in information and communications technology. The 2007 HMP Jumpstart signaled the propulsion of microbiome research. Early human microbiome research started with simple studies on culture methods. However, it later showed massive progression with an explosion of data and developments in metagenomics. From 2012 onwards, national level studies started taking place in Korea, France, Australia, Japan, and other countries, in addition to existing projects led by the US and the EU (Table 1)\textsuperscript{34}). Research areas expanded to include respiratory organs, skin, oral cavity, and female...
reproductive organs.

1. International Human Microbiome Consortium

The International Human Microbiome Consortium (IHMC) was organized in order to analyze and share genetic information from specific disease causing microbes globally. There are currently nine member states, and Korea joined in 2011. Major research programs include the HMP and National Microbiome Initiative (NMI) in the US, MetaGenoPolis (MGP) and International Human Microbiome Standards (IHMS) led by France, the Korean Microbiome Diversity Using Korean Twin Cohort Project by Korea, the Australian Jumpstart HMP supported by the Australian government, Canadian Microbiome Initiative (CMI) supported by the Canadian government and The Metagenomics of the Human Intestinal Tract (MetaHIT) funded by the European Commission.

2. Human Microbiome Project

The US implemented HMP as a 5-year research plan at the federal level, supervised by the National Institutes of Health (NIH), in order to identify and characterize the microbes that affect human health and disease. HMP started with the goal of completing an exhaustive map after analyzing the genome base pairs of approximately 3,000 species of microbes inhabiting five parts of our body, namely the oral cavity, skin, vagina, intestinal tract, and lungs. Although the exact causative relationships are

| Table 1. Major International Microbiome Projects |
|-----------------------------------------------|
| Country | Project title | Duration | Objective |
|---------|---------------|----------|-----------|
| International | International Human Microbiome Consortium (IHMC) | 2010 ~ | Set principles and policies to study microbiome. Mediate projects to generate a comprehensive data resource. |
| EU | Metagenomics of the Human Intestinal Tract (MetaHIT) | 2008 ~ 2011 | Construct large catalog of intestinal microbiome and genes of IBD and obesity. |
| | International Human Microbiome Standards (IHMS) | 2011 ~ 2015 | Establish a standard protocol for human microbiome studies. |
| USA | NIH Human Microbiome Project (HMP) | 2008 ~ 2015 | Generate resources of the human microbiome. Diagnosis and treatment through a correlation between diseases and microbiome. |
| | Home Microbiome Project | 2012 ~ | Analyze bacterial community in living space. |
| | Data Analysis and Coordination Center (DACC) | 2008 ~ 2013 | Assist in standard of data pipeline. |
| | Hospital Microbiome Project | 2012 ~ 2014 | Analyze bacterial community in hospital, and find correlation between the bacteria and patient. |
| | Microbiome Quality Control Project (MBQC) | 2013 ~ | Establish analytic pipeline of human fecal microbiome. |
| | American Gut Project | 2013 ~ | Study of bacterial diversity of the public and construct data resource. |
| Japan | Japanese Consortium for Human Microbiome (JCHM) | 2014 ~ | Study of Japanese specific intestinal microbiome and disease biomarker. |
| Canada | Canadian Microbiome Initiative (CMI) | 2009 ~ | Support seven research teams to conduct a prospective study of diseases by modeling and mapping microbial diversity. |
| France | MicroObes | 2008 ~ 2010 | Research on obesity and human intestinal microbiome. |
| | MetaGenoPolis (MGP) | 2012 ~ 2019 | Find the correlation of non-infectious diseases and intestinal microbiome. |
| Korea | Korean Microbiome Diversity Using Korean Twin Cohort Project | 2010 ~ 2015 | Study on human diseases related microbiome using twin cohort and characterize Korean specific microbiome. |
| Australia | The Australian Jumpstart Human Microbiome Project | 2009 | Analyze metagenome of intestinal microbes and Australian specific microbiome. |

Adapted from the article of Choi et al. (Allergy Asthma Respir Dis 4: 311-320, 2016) in accordance with the Creative Commons Attribution Non-Commercial (CC BY-NC 4.0) license. IBD: inflammatory bowel disease.
yet to be revealed, the gut microbiota composition of people with various diseases differs greatly from that of healthy people, and gut microbiota differences between people with autism, mental illnesses, and Alzheimer’s and healthy people have been published. The ultimate goal of the HMP is to reveal the association between disease and human microbiome changes, and to identify correlations between the human microbiome and disease and health.

3. Metagenomics of the Human Intestinal Tract

The MetaHIT project, that started in 2008 and was led by French researchers, was active until 2012 with participation from 15 institutions in 8 countries (France, Denmark, Germany, Spain, Italy, the Netherlands, the UK, and China). The goal of MetaHIT was to identify the role of the gut microbiome linked with IBD and obesity, and to analyze their correlations. As a result, a reference catalog of microbial genes comprising the gut microbiome was constructed, and a method capable of analyzing gene frequency differences from individual distribution of microbiota was developed.

4. MetaGenoPolis

The MGP was founded in order to investigate influences of gut microbiota on noninfectious diseases, and to examine how the gut microbiome affects human health and infectious diseases through interactions between microbes. Human gut microbiome analysis and genome analysis were conducted, bioinformatics tools for sample analysis were developed, and databases were constructed from results of the analysis.

5. Human MetaGenome Consortium Japan

In 2004, Japanese researchers discovered a link between gut microbiota and the brain by observing a twofold increase in the release of stress hormone in a mouse after removal of its gut microbiota. This study was the starting point that led to the establishment of the Human MetaGenome Consortium Japan (HMGJ). The project aims to identify distinct gut microbiome characteristics in the Japanese population, and construct a Japanese specific database. It is a large-scale microbiome project which collects, shares, integrates, and manages human and animal microbiome research data in Japan, and corporate and government research institutes are its participants.

6. Beijing Genome Institute

China has been conducting a joint microbiome research project with Europe and Japan. With the goal of completing the 10,000 Microbial Genome Project, Beijing Genome Institute (BGI) has been conducting research on various microbes with diverse applications in agriculture and livestock industry, bioenergy, the environment, and medicine since 2009. It currently plans to conduct the Chinese Microbiome Project.

Conclusion

Microbiome research studies the diverse microbes inside the human body, and microbiome industry is gaining interest as a new growth driver in biotechnology, which is an essential component of the Fourth Industrial Revolution. Expectations are high about the possibility of novel diagnostic strategies and drug development based on the microbiome, as research reveals the microbiome’s impact on a number of diseases including bowel diseases, atopic dermatitis, obesity, metabolic diseases, and depression. While microbiome research has been active globally, many studies still simply settle for evidence that microbial imbalance in the body affects a disease. Although microbiome research is yet to be fully established in Korea, the government has been investing on human microbiome studies since 2016, especially on studies, which can improve therapeutic strategies or health. Predicted marketability is high for microbiome-based medicine, as it can be applied to invent cures not only for autoimmune diseases such as rheumatoid arthritis, but also for gastrointestinal diseases such as Crohn’s disease and IBD, diabetes, and so on.

A healthy oral cavity is the home for many ordinary bacteria, fungi, and viruses. Over 600 species are estimated to inhabit the oral cavity, some of which are pathogenic while others are mutualistic. When oral and gut microbiota of a healthy group and a group with metabolic disease were compared in a twin study of oral microbiota in 228 people, the two groups showed a significant difference in
microbiota diversity\(^43\). *Neisseria* and *Granulicatella* were more visible in the oral cavity of the group with metabolic diseases, and quantitative investigation using quantitative polymerase chain reaction obtained the same result\(^43\). These oral microbes associated with metabolic diseases are expected to serve as biomarkers for predicting drug response, and assist in research about their pathogenesis mechanisms. In 2016 a survey on male and female adults, periodontal disease was found to be significantly correlated with other diseases including angina, cerebral infarction, and cardiovascular diseases such as myocardial infarction, rheumatoid arthritis, diabetes, and osteoporosis\(^44-49\).

*Porphyromonas gingivalis*, which causes periodontal diseases, can also enter the bloodstream and cause the formation of atheroma\(^50,51\). Reports also indicate that it can activate platelets and induce platelet-leukocyte aggregates, causing a prothrombotic state\(^50,51\). Prevalence of bacteremia and endotoxemia were higher in patients with periodontitis than patients with gingivitis or healthy people. *P. gingivalis*, *Actinobacillus actinomycetemcomitans* and other periodontal pathogens were found in atherosclerotic tissue biopsies from patients with periodontal disease\(^52\).

Most dental diseases such as gingivitis, dental caries, halitosis, and stomatitis can be attributed to the activity of harmful oral bacteria, and recent claims about their effect on systemic health are drawing attention to the need to maintain a healthy oral microbial ecosystem. Therefore, a need for research methods to analyze oral microbiome and a plan for long-term research is increasing.

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**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.

**ORCID**

Sung-Kyun Chung, https://orcid.org/0000-0002-4022-1620
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