Microbial Diversity: Relevance and Relationship Between Environmental Conservation And Human Health

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ABSTRACT

This work presents bibliographic data on the role and function of microbial diversity. The increasing use of probiotics and prebiotics foods has led to the studies on their actual functions in the human body. It is known that in the environment, microorganisms are extremely important in recycling of nutrients, balance of trophic chains, vital physiological activities in the plants and animals, as well as the conservation of natural habitats. In human food, these microscopic organisms contribute from flavoring products to the synthesis of antimicrobial substances and vitamins essential to living beings.

Key words: Bacteria, nutraceutical, collective health, review

INTRODUCTION

Environmental and health issues, along with the increase in food production, have been topics of investments in the area of scientific research. Food production with quality and quantity to meet the world’s population is a matter related to public health. The defense of biodiversity stimulates the production of foods that provide benefits for the consumers and simultaneously benefit environmental sustainability.

Consumption of processed products with low nutritional value and high levels of trans fatty acids is frequent in the population. In Brazil, ten steps to a healthy diet were included in the National Plan for the promotion of proper nutrition and healthy weight, which aimed to improve people’s knowledge about health and encourages the practice of physical exercises. In these ten steps, it is possible to check the improvement in the quality of life of the consumer, so they facilitate and improve the eating habits of the intestinal microbiota, and therefore, a person’s health (Vinholes et al. 2009).

The human diet is a key indicator of quality of life and affects individuals in different ways, because of the importance of protein, vitamins, minerals and nutrients that are needed for the perfect functioning of the body. The cereals have similar percentages of protein, lipid, fiber, ash and carbohydrate. Cereals are foods of plant origin, consisting of grains and largely consumed by the people worldwide. The name ‘cereal’ derives from the \textit{Ceres}, the Greek goddess of agriculture and harvest. The main cultivated cereals are rice, wheat, corn, oats, rye, barley and triticale. They are widely consumed for being part of the food habits of many people; for ease of culture, conservation, transportation and income, for being

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low cost and good nutritional value and for the
great variety of forms of use (Coppens 2005).
Rice is one of the most cultivated cereals in the
world for human consumption and raw material
for the food industry. There are increasing trends
of these industries to seek rice, mainly organic and
whole grain rice, due to health issues and the
media, which has an interest in the cultural revival
of a power based on wholegrain products (Ley et
al. 2005). Rice, despite not being a food rich in
vitamins, has a differential to be easily assimilated
by the body. Its main component is starch, which
provides energy, serving as fuel for the
functioning of the body. The protein percentage
varies around 3 to 12% depending on the type of
rice (Ibge 2013).
Practitioners of macrobiotic and vegetarian diets
seek to use rationally, using the rice in order to
promote a complementary action of one food over
another. In nature, the microbial communities
facilitate the flow of nutrients and energy from the
earth, but little is known about the interaction, and
their diversity is still underestimated. Their
patterns of succession and how their spatial
distribution also occurs are poorly studied
(Barbosa et al. 2010). Therefore, functional
biodiversity in agroecosystems is key to the
ecological sustainability of production and
microorganisms are essential to this process
(Johnson et al. 2003).
Bacteria comprise a large microbial community,
consisting of species considered pathogenic to
humans, animals and plants, or even beneficial
species that interact with other organisms.
Therefore, the importance of the study of
microbial diversity is at stake.

METHODS

The bibliographic search was performed through a
systematic search covering the 1900s and up to
2013. The terms “soil bacteria”, “agroecosystem”
and “nutraceutical” were utilized to search
databases like SciELO and Portal Capes. These
databases fulfill the minimum criteria search to
conduct a systematic review, in accordance with
the literature. The articles written in languages
other than English, Spanish or Portuguese were
excluded. Then the full text of selected articles and
cited references were analyzed, which were
selected in additional studies that belonged to the
subject. The initial search identified 68 articles,
which were used for the preparation of this review
article

MICROBIAL DIVERSITY
In preserved ecosystems
Microorganisms represent the richest repertoire in
chemistry and molecular diversity in nature,
providing the basis for ecological processes such
as biogeochemical cycles and food chains, as well
as maintaining vital relationships among
themselves and with superior organisms (Hunter-
Cevera 1998). The diversity of microorganisms is
as vast as it is unknown. So far, between 0.1 and
10% of microbial species are known, depending
on the habitat studied. In an agroecosystem, the
variation in microbial diversity throughout the
seasons is still not well understood, since in each
season, one microbial community seems to occur
more dominantly, accompanied by other less
abundant that often are below the level of
detection using the current methods of evaluation
(Torsvik and Ovreas 2002). The diversity of
microorganisms is critical to the functioning of
the ecosystem, because there is the need to maintain
ecological processes such as decomposition of
organic matter, nutrient cycling, soil aggregation
and controlling pathogens within the ecosystem
(Kennedy 1999). The functional diversity is very
important in ecological assessments of
microorganisms within the ecosystem, mainly
because little is known about the relationship
between the structural and functional diversity of
these microorganisms. There is, however, a
consensus that microbial diversity is directly
related to ecosystem stability (Yamanaka et al.
2003).

In agroecosystems
Modern agriculture is characterized by a constant
search for increased crop yields through the use of
mechanization, irrigation, chemical fertilization
and pesticide application, coupled with the
improvement of plant genotypes. However, the
environmental impact caused by the intensification
of farming did not always receive the necessary
attention. The lack of detailed knowledge about the
ecosystem and/or inadequate planning for land
use led to a framework of intense environmental
degradation, with loss of non-renewable resources
and biodiversity not only in Brazil but in other
countries (Eussen 1997). The observation in the
increase of degraded agricultural areas has led to a
change in the concept of land use, based on a
holistic view of the agricultural process, where natural resources (soil, water and biodiversity) are exploited more sustainably. The diversity of microorganisms as an indicator of the quality of agroecosystems has been widely debated, especially in the last decade, with the advent of molecular biology techniques that have favored the evaluation of microorganisms in environmental samples (Coutinho et al. 1999; Tiedje et al. 2001; Unicomb et al. 2005). The main argument in favor of such environmental feature is the fact that the microbial diversity naturally remains unchanged throughout the year (Dickens and Anderson 2001). Despite the continued use of land for agricultural practices, the soil may be rich in the biological point of view, since many microorganisms found in this environment are considered important in the biological control of diseases and pests of agriculture (Andreoti 2009). Thus, microbial processes play a fundamental importance in the functioning of production systems, performing tasks directly related to their productivity and sustainability (Bonkowski and Roy 2005). Rice is a plant of the genus *Oryza* of the grass family. *O. sativa* (Asian rice) and *O. glaberima* (African rice) are the two most cultivated species. The domestication of rice happened about 10,000 years ago in Asia. In Brazil, the plant was introduced by the Portuguese (Heinrichs 1997; Hunter-Cevera 1998). Rice is still considered the main component of the world's staple diet, although the per capita consumption in Brazil has decreased by 40.5%, according to the study by the Brazilian Institute of Geography and Statistics (FAO) in 2013. However, it is noteworthy that the production of rice in 2013 was 12,816 million tons. Rice cultivation in Brazil is primarily in irrigated mode. The largest producing state of Brazil is Rio Grande do Sul with 1,0666 million hectares, representing 44.5% of the national area and accounting for 66.5% of Brazilian production (Panizzon et al. 2013).

Microbiological monitoring of water and soil depends on the quality of water entering the crops. Part of the diversity of Gram-positive bacteria found in rice ecosystems belong to the groups *Bacillus*, *Micrococcus*, *Staphylococcus*, *Corynebacterium* and *Lactobacillus*, among with a variety of Gram-negative species belonging to the families *Enterobacteriaceae*, and *Pseudomonadaceae* *Aeromonadaceae*, *Vibrionaceae*, *Brucellaceae*, *Burkholderiaceae*, *Xanthomonadaceae* and others (Reche and Fiuza 2005). Among the Gram-negative bacteria, the following species have been identified by the same authors in rice cultivation waters: *Escherichia coli*, *Enterobacter* sp. *Citrobacter freundii*, *Klebsiella pneumoniae*, *K. oxytoca*, *Proteus mirabilis*, *P. vulgaris*, *P. pomerii*, *Citrobacter* spp, *Enterobacter cloacae*, *Providencia rettgeri*, *Stenotrophomonas maltophilia*, *Ochrobactrum anthropi*, *Chromobacterium violaceum*, *Yersinia enterocolitica*, *Pseudomonas aeruginosa*, *P. fluorescens* and *Burkholderia cepacia*.

**In Human microbiota**

The human body is populated by a large number of microorganisms, being distributed in various tissues and organs. It is believed that the abundance of microorganisms is greater than the number of cells in the body and is directly related to the immune system and better utilization of feed (Brandt et al. 2006). The digestive system consists of the alimentary tract associated with organs that partially perform digestion, mechanically and chemically. The purpose of the digestive system is the breakdown of food into smaller particles in order to be absorbed in human body.

The gastrointestinal tract of mammals maintains a highly diverse microbial population that plays an important role in nutrition metabolism, protection against pathogens, and the development of the immune system. It is estimated that at least 1000 different bacterial species coexist the human intestinal tract. Although often considered as pathogens, the majority of microorganisms in intestinal tract have beneficial effects. They play multiple roles in the human host, as they are directly involved in the synthesis of vitamins and cofactors, help to degrade complex lipids and polysaccharides and also have detoxifying action (Kennedy 1999; Douglas- Escobar et al. 2006). It is known that bacteria intensely populate the most part of the enteric tract. However, the stomach and the small intestine have fewer microorganisms due to the presence of hydrochloric acid (Bedani et al. 2009). The large intestine has numerous bacteria, especially *Lactobacillus* sp., *Escherichia coli*, *Klebsiella* sp. and *Proteus* sp. that assist in the synthesis of degradation of the food (Camargo et al. 2011; Odonkor and Ampofo 2013).

The intestinal microbiota is a dynamic mixture of microorganisms, whose composition varies among the gastrointestinal tract (GIT) and between the
mucosa and the intestinal lumen. The microbiota develops all the time, due to the interaction of genetic factors, contact with the environment, diet and disease, thus explaining the fact that each individual presents a unique microbiota (Bedani et al. 2009). The genotype of the individual may also be a factor that influences the bacterial community in the gut, following a hypothesis that in the gut there are sites of specific adhesion to some bacteria in the host, selecting the most important. These adhesion sites are probably genetically predetermined (Brandt et al. 2006). Figure 1 represents didactically the factors that can influence the intestinal microbiota.

![Figure 1](image)

**Figure 1** - Factors influencing the bacterial community in the human gut.

The highest concentration of microorganisms and metabolic activity is found in the large intestine. Starting from the ileum the concentration of bacteria increases gradually, reaching $10^{11}$ to $10^{12}$ CFU/g in the colon. The adult and stable microbiota are composed of autochthonous species (permanent members) and allochthonous (transient members that are acquired from an external source). The microbiota are diverse, comprising 400-1000 species, of which over 60% are not cultivable outside the intestine (Moreira 2012). The intestinal microbiota plays different roles that are important for the host such as they exert a trophic effect on the intestinal epithelium, favoring the development of microvilli, which in turn promotes the absorption of the nutrients (Moreira 2012). The diet is a major determinant for the persistence of certain bacteria in the gastrointestinal tract, because the diet provides nutrients, not only for the host, but also for bacteria that are in it. Components from indigestible food serve as carbon and energy source for the human gut bacteria (Blaut and Clavel 2007).

The intestinal biota is increasingly considered a symbiotic partner for the maintenance of health. Over the years, the intestinal microbiota has become recognized in the development of the immune system. The cells of the intestinal mucosa and microbiota (enterocytes, dendritic cells, lymphocytes, macrophages) regulate the production of several cytokines and chemokines. The homeostasis of the intestinal tract is dependent on characteristics of the host (age, gender, genetics, etc) and environmental conditions (stress, drugs, gastrointestinal surgery, infectious and toxic agents). Changes in the gut microorganisms can be reversed by diet and weight loss. Qualitative and quantitative changes in the consumption of specific food components (fatty acids, carbohydrates, micronutrients) have effects not only on the composition of intestinal microbiota, but can modulate the expression of genes in the host tissues such as the liver, adipose tissue, intestine, and muscle. This in turn can lead to the development or decrease of fat mass and metabolic disorders associated with the intestinal barrier and systemic immunity. These intestinal microbiota are composed of various bacterial groups. Therefore, it is essential to keep them balanced to preserve human health.

Some bacteria have been used as a probiotic, such as species of the genera *Lactobacillus* and *Bifidobacterium*, which help in the restoration of the human intestinal microbiota, when the intestinal biota is drastically reduced and is vulnerable to attack by the pathogens (Santos and Cançado 2009).

**Human nutraceutical**

The evolution of nutritional science made it possible to know a little more about the food composition and action of these substances in the body. Studies have been carried out on the phytochemical constituents of the nutraceuticals in the food in order to study their health benefits. Nutraceuticals are dietary supplements that contain a concentrated bioactive compound in the food. They are used for improving the health that could be obtained from the food (Zeisel 1999).

The Brazilian Health Surveillance Agency of the Ministry of Health in the ordinance n° 398, April 30, 1999 defined the functional food as the ingredient that claimed functional or health properties. It may, in addition to basic nutritional

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functions, in the case of a nutrient, and produce metabolic or physiological effects and or health benefits, and should be safe for consumption without medical supervision (Vigilância Sanitária 1999). The importance of the adequacy of food with respect to microorganisms in order to reduce the cost, increase quality regarding the taste and nutrition must be kept in mind. They produce a more flavorful food, with less cost and less waste of chemicals, which is of great significance for the food industry (Ley et al. 2005; Panizzon et al. 2012).

**Prebiotic and probiotic microorganism**

The gastrointestinal tract is colonized by the microorganisms soon after the birth of the individual and also after certain life situations, such as the consumption of antibiotics, resulting in the reduction of microorganisms in the gut, which makes it important to understand when this type of food should be used. The enterobacteria and some species of *Clostridium* spp. are harmful to human health. The gut biota balance is maintained by the interrelations between the bacteria themselves and the host. Accordingly, the probiotics increase the number of microorganisms in the host; in addition, they induce pro-inflammatory, anti-inflammatory and anti-carcinogenic actions. The majority of probiotic microorganisms are lactic acid bacteria, which are Gram-positive and microaerophilic (Rodrigues and Barroso 2011). Probiotics are dietary supplements added to the food in adequate quantities influencing their nutritional and therapeutic value. They are represented by cultures of live microorganisms (beneficial bacteria and yeasts), which contribute to improve the flavor of the final product and produce antimicrobial substances (Santos 2009). Prebiotics are food ingredients that are primarily substrates for the growth of the intestinal microorganisms, not digested in the small intestine that, upon reaching the large intestine, are selectively metabolized by a limited number of bacteria, called beneficial, which alter the microbiota of the colon generating a healthy bacterial microbiota (Raizel et al. 2011).

A healthy diet by adding probiotics and prebiotics in food benefits the people. In today’s society, with the increasing population life expectancy and drug costs, nutraceutical should unify scientific knowledge about the public health and nutrition suitable to these new challenges by promoting health and not just curing diseases. The healthy microbiota promotes well-being and with the addition of probiotics, it is possible that they could produce antagonistic effects against unwanted microorganisms. Therefore, knowledge of the gut biota leads to the development of numerous public health strategies (Saad 2006). The importance of probiotic approach to the management of obesity in humans is supported by some intervention studies in humans so far, but the experimental data obtained with the compounds help to elucidate potential new molecular targets associated with the diet and the existing bacteria (Delzenne et al. 2011).

Recent studies involving rice bran and its oil have shown that they possess prebiotic potential, aiding in the growth of important microorganisms in the intestine, such as those of the genus *Lactobacillus* that are highly effective in controlling the intestinal infections by *Salmonella* and are catalysts of non-digestible molecules (Kumar et al. 2012; Tamura et al. 2012). Changes in the bacterial composition of obese versus lean people relate to bacterial phyla such as, for example, a reduction in *Bifidobacterium* spp. or an increase of *Staphylococcus aureus* (Neyrink and Delzenne 2010).

**Pathogenic microorganisms**

The members of the normal microbiota can cause diseases under certain circumstances. Since they have a non-invasive way of life defined by limitations of the medium, unless they are held they can become pathogenic. Population levels of microorganisms are determined by the exogenous and endogenous multifactorial processes (Griffiths 2001). The bacteria of the intestinal tract have heterogeneous distribution. The colonization of the intestinal tract depends on the ability of bacterial adhesion. There are bacteria on the adhesion sites on the intestinal mucosa, which need not to be periodically reintroduced. However, there is the native biota that is external to the gut ecosystem, thus is transient. The microbiota has the following functions: antibacterial, immunomodulatory and metabolic. Antibacterial prevents the establishment of pathogenic bacteria. Immunomodulatory activity helps the immune system and metabolic function contributes to facilitate the nutrition (Brandt et al. 2006). It is important to highlight the impact that pathogenic bacteria can cause in the public health issue, resulting serious intestinal diseases such as diarrhea - considered as the most common disease.
caused by viruses and bacteria and one of the diseases that affects large no of children in the world. Hence, it is important to know the bacteria that may possibly compromise the gut and the human organism as a whole (Clotildes 2007). The use of antibiotics in the rats can increase intestinal microbiota associating to some changes that affect the acquisition of energy from compounds in the diet and how it is spent and stored (Ley et al. 2005). Salmonella is represented by more than 40 serogroups and 2000 serotypes and may be classified as typhoid and non-typhoid. This genre is usually associated with food. Salmonella enteritidis (Fig. 2B) is one of the serotypes most widely distributed in the world and one of the major contaminants in food, usually beef, pork, poultry and eggs. This bacterium usually causes fever, abdominal cramps and diarrhea, which can present blood clots. A study by the Center for Epidemiological Surveillance - SES / SP showed that from 1999 to 2007, S. enteritidis was responsible for 42.3% of outbreaks of diarrhea, showing the attention in public health that this bacterium should be given (Kirk et al. 2004; Unicomb et al. 2005; Paiao et al. 2013).

Escherichia coli (EPEC) (Fig. 2C) causes gastroenteritis in almost all age groups. It is similar to Shigella sp (Fig. 2D) because it penetrates directly into the intestinal epithelium where it can multiply, causing dysentery. It can be transmitted by the consumption of water and many foods such as milk and milk products. The importance of EPEC as a cause of diarrhea has declined since the 1960s, but is the primary infectious agent in children in developing countries, including South America, Africa and Asia. EPEC outbreaks are sporadic, emerging in places where sanitary conditions are poor (Silva et al. 2001; Maltick et al. 2010). Other bacteria that are not involved directly with the human gut, but are cause of worry in the matter of public health, are related to urogenital infections, such as Proteus vulgaris (Fig 2A), which inhabits the human gut, but causes urinary tract infections and other complications (Rodrigues and Barroso 2011).

Having a prebiotic diet can help treating the intestine infections, such as those caused by the bacteria of the genus Salmonella because these prebiotic molecules serve as substrate for the growth of bacteria, which has the potential to eradicate other pathogenic bacteria such as Lactobacillus (Kumar et al. 2012).

Microbial diseases of the large intestine are second only to the respiratory system diseases. Pathogens are able to cross the digestive system and extend to organs, causing numerous diseases, e.g., gastroenteritis caused by Salmonella and rotavirus. Bacillus cereus is common in the soils and vegetables. Rice has vast abundance of this bacterium. It is generally harmless, but when found in the foods, can cause illnesses such as gastroenteritis. The rice, for being a plant that is cultivated in water, has the risk of receiving numerous microorganisms. Water is characterized by low nutrients. Therefore, bacteria tend to grow on standing surfaces in particular materials, as is the case with rice (Pomeroy 1974a; Pomeroy et al. 2007b; Andreoti et al. 2009). Therefore, paddy fields are important for local biodiversity conservation because they support a rich biodiversity and high productivity feature. Rice is one of the most important cereal crops in the world. Therefore, the conservation of biodiversity in agriculture is a challenge of great importance. Several studies have demonstrated the contribution of ecosystems such as rice, providing habitats for creation of numerous microorganisms (Hofman et al. 2003; Maltick et al. 2010). The microorganisms in the biosphere perform important functions, for example, the influence on biogeochemical processes. In aquatic environments, there is an important chain of interactions that affect the elements involved in the environment (Comte et al. 2006). It is noteworthy that quality evaluations in health should always aim at the welfare of the patient, which is always the focus of the studies. Therefore, monitoring allows detecting the faults and correcting them in

![Figure 2 - Photomicrographs of bacteria (100x) for representing their morphology.](image-url)
order to not compromise the product and the consumer (Abrantes et al. 2007). It is possible to relate the bacteria of rice with their intake and activity within the human body. Microorganisms can generate many by-products through fermentation, which may even have commercial value and are easily produced. The cost/price margin is small because they can be easily replaced by cheap products and chemicals; however, they are naturally formed in the body. The development of non-dairy probiotic products is a major challenge in the food industry because many are lost during the processing and storage of the product. Cereals such as rice have been widely studied and can be used as fermentable substrates that favor the growth of probiotic microorganisms (Pomeroy et al. 2007b; Oliveira and Jurkiewicz 2009). Microorganisms have a high degree of specificity, acting separately in chemical reactions. Therefore, with the development of biotechnology, it could be possible to manufacture the food and perform transformations that can affect people's lives.

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