

Name MG Lab Gender Age Sample code MPC00174 Registration date 27-06-2023 Sample date 18-07-2023 Sample received date 26-06-2023

Summary

People develop an enterotype that will remain with them for most of their lives in the first three years of their life. These bacterial groups thrive on different food sources but fulfil a similar function. The enterotype mostly provides insight into a person s diet.

Enterotype 1 Bacteroides This enterotype is characterized by a dominance of the genus Bacteroides. Bacteroides live off the carbohydrates of the mucus layer that your gut produces. The genus is often seen in people with a diet rich in meat. However, even people with a plant based diet can have a Bacteroides enterotype, if they do not have Prevotella in their microbiome

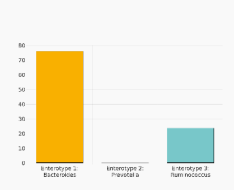
Enterotype 2

Prevotella Characterized by a dominance of the genus Prevotella. Prevotella lives off plant-based fibres and is therefore often found in people with plant based diets Unlike Bacteroides, Prevotella is not able to digest the carbo hydrates that are pro vided by the mucus layer of the gut.

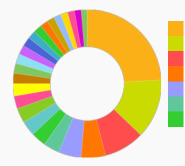
Enterotype 3: Ruminococcus

Characterized by a dominance of the genus Ruminococ cus. They are often found in people with a higher intake of resistant starches. This enterotype is less clearly defined and given certain conditi ons (specific nutrition) this enterotype can transition to entero type 1 or 2



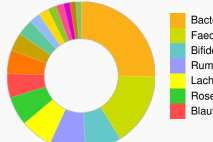


Genus level microbiome of people with your enterotype



Bacteroides (18.8%) Faecalibacterium (10%) Blautia (6.9%) Eubacterium (4.1%) Ruminococcus (3.8%) Alistipes (2.7%) Roseburia (2.6%)

Your genus level microbiome



Bacteroides (21.3%) Faecalibacterium (13.6%) Bifidobacterium (6.7%) Ruminococcus (6.7%) Lachnospiraceae (5.9%) Roseburia (5.3%) Blautia (4.4%)

Species balance

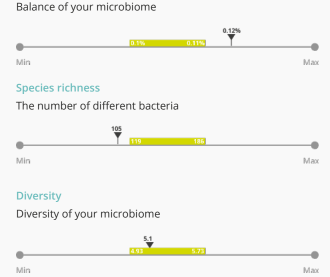
The species evenness (Pielou index) provides an impression of evenness in the distribution of your microbiome. A high species evenness is a measure of a very uniform distribution of your microbiome. A low species evenness, on the other hand, is related to an uneven distribution of your microbiome.

The species richness is a measure for the number of different bacteria species present in your gut. The diversity (Shannon index) combines these two parameters and is a measure of how diverse your microbiome is.

For example, a low diversity (Shannon index) can be the result of a low number of different bacteria (species richness), but this can also be the result of several bacteria that dominate the gut microbiome, which is reflected by a low species evenness.

Different diversity scores with reference values

Species evenness





Clinically Relevant Parameter

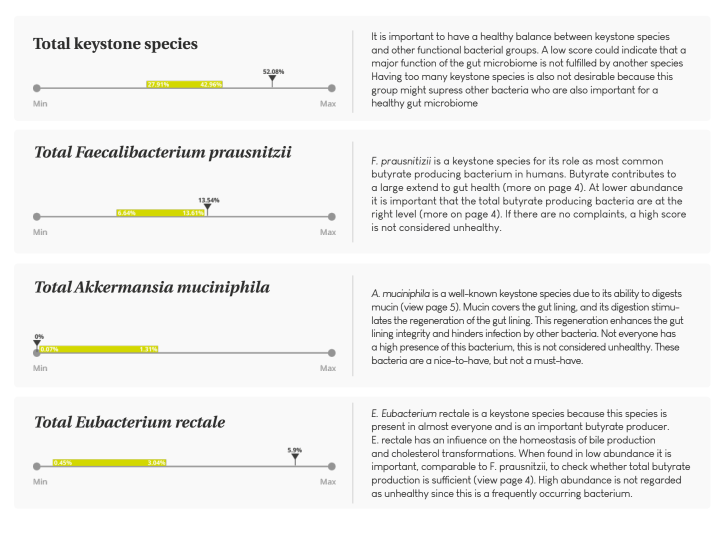


The microbiome in this report is compared with healthy individuals in the MyMicroZoo database. The upper and lower bounds of the reference bars of the respective bacteria are based on the bacterial abundance in 50% of the healthy individuals. The microbiome is compared with the microbiome of individuals with the same enterotype and who are over 18 years old, non-smokers, do not suffer from gut related complaints or other illness and have not used medication or antibiotics in the past 6 months.

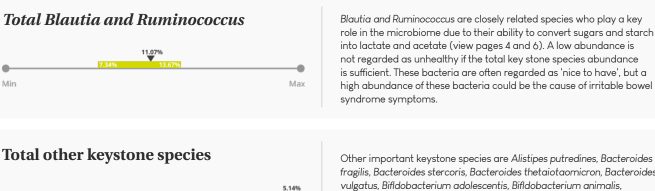
Please note: Results outside of the green reference values are not necessarily considered unhealthy if there are no complaints. We recommend speaking with a healthcare professional when complaints are present. A healthcare professional can help to provide personalized advice based on your gut microbiome.

Keystone species

As the name suggests, keystone species fulfil a key role in the microbiota. Even though these species are often found in low abundance, they have a great effect on the gut microbiome. There is a great effect on other bacteria if keystone species are lost. This is because keystone species often play an essential role in the food chain or in communication between bacteria. Often, keystone species fulfil one or more roles that are not easily transferred to different species







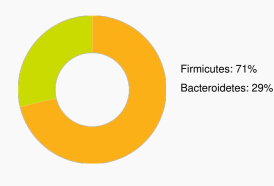
0.48% 2.47% 5.14% Min Max

Other important keystone species are Alistipes putredines, Bacteroides fragilis, Bacteroides stercoris, Bacteroides thetaiotaomicron, Bacteroides vulgatus, Bifldobacterium adolescentis, Bifldobacterium animalis, Bifldobacterium longum, Christensenella minuta, Oscillospira, Roseburia. Members of these species are seldom found in larger quantities than a few percent.

The parameters above provide a general overview. When deviations are seen, more detailed information can be found by looking at the present of individual presence per genus (view page 9 'Genus level analysis' or view the scores per species in the online results)

F/B Ratio

Firmicutes/Bacteroidetes ratio (F/B ratio)

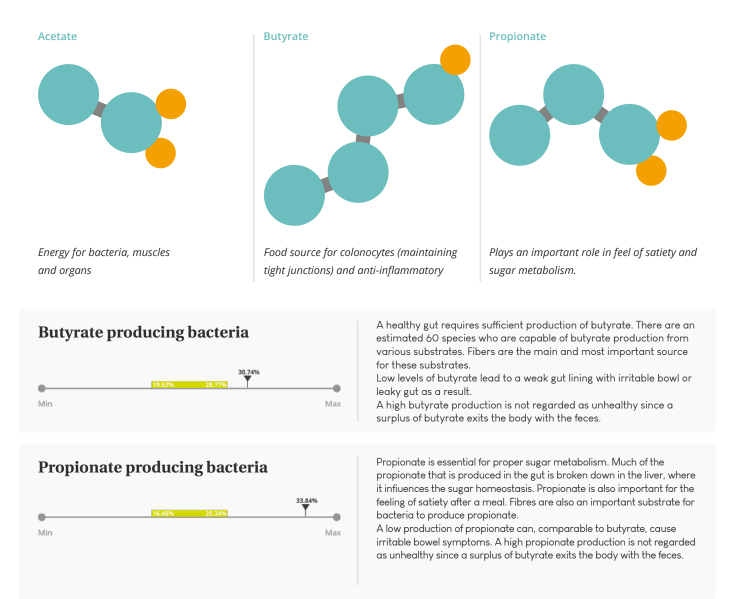


Firmicutes and Bacteroides are two phyla who together can account for 90% of the microbiome. The F/B ratio varies wildly between individuals and is very dependent on the diet. However, it is known that Firmicutes is more often seen in the developed world and is also correlated with obesity. Bacteroidetes is more often found in the developing countries with a more traditional diet and is correlated with irritable bowel symptoms.



Short chain fatty acids (SCFAs)

The short chain fatty acids: acetate, propionate and butyrate are the most important substances produced by the microbiome. Acetate is a food source for many other bacteria of the microbiome and is often provided to the bloodstream to be used as an energy source for our organs and muscles. Propionate plays a major part in the feeling of satiety and in the sugar metabolism. A large portion of propionate is processed in the liver. Butyrate is a direct food source for the cell that make up our gut lining; the colonocytes. Healthy colonocytes provide the gut lining of structural integrity which inhibits 'leaky gut'. This is done by reenforcing the tight junctions. Tight junctions are the strong connections between colonocytes. When colonocytes do not have the necessary food sources (short chain fatty acids) the tight junctions will weaken creating 'leaks'. This causes 'leaky gut' and is often associated with infiammation. The unhealthy colonocytes also contribute to an increased rise in (colon)cancer due to the worsened cell division.

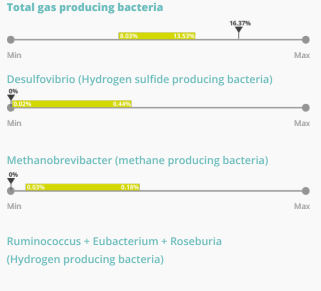


Footnote

Many bacteria can produce different short chain fatty acids. Whether a bacteria produces acetate, propionate or butyrate is dependent on the substrates available to the bacteria. Butyrate and propionate production can be estimated based on the butyrate and propionate producing bacteria that are found in the gut microbiome. It can be said for both metabolites that a high production is favourable. However, it is not possible to estimate acetate production based on the amount of acetate producing bacteria found in the gut microbiome. Which bacteria actually produce acetate is much more dependent on the substrate intake (diet) than for butyrate or propionate.



Gas producing bacteria



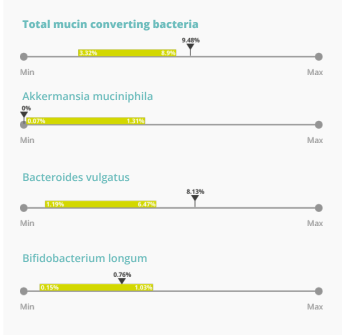


Certain bacteria in the gut will produce different types of gas when they are fermenting food sources. Food sources like certain sugars or non-digestible polysaccharides (like starch and cellulose) are not digested in the small intestine but are fermented in the large intestine by gas producing bacteria. The gasses that these bacteria produce are mainly carbon dioxide (CO2), hydrogen (H2), methane (CH4) and hydrogen sulfide (H2S) Gas production in the gut is a natural process which provides the gut and body with energy. Most of the produced gas is used by bacteria in other processes. A smaller part of the produced gas exits the body as fiatulence or is exhaled after it is transferred in the blood to the lungs.

A high abundance of gas producing bacteria can cause different intestinal complaints, such as bloating or excess fiatulence. This can be reduced by decreasing the intake of the food sources of the gas producing bacteria. A low-FODMAP diet (fermentable oligosaccharides, disaccharides, monosaccharides and polyols) can help to decrease the abundance of gas producing bacteria. Important food groups that are eliminated are beans, but also sugar from milk products (lactate) or sugars from fruit (fructose) are excluded in a low-FODMAP diet.

A low abundance of gas producing bacteria does is not regarded as unhealthy for the (gut) health.

Mucin converting bacteria



Mucus is a slimy layer that is produced by the lining of our gut. The building blocks of this mucus layer are mucins. Mucins are important for our gut health because they function as lubrication, hydration for the gut lining, protection against pathogenic microbes and as a food source for mucin converting bacteria. Mucin converting bacteria are important because when they digest the mucins, they also stimulate the renewal of the gut lining and stimulate the production of new mucins. *Akkermansia muciniphila* is an example of a mucin converting bacteria that converts mucins into products which are used by other bacteria, such as Bacteroides, as a food source while also stimulating the renewal of the gut lining and the production of mucins.

A low abundance of mucin converting bacteria is often seen with a reduced gut lining renewal and reduced mucin production. This is regarded as unfavourable due to the many functions of mucins. A high abundance of mucin converting bacteria is not regarded as unhealthy for the (gut) health.



Sulphate reducing bacteria

Total Sulfate Reducing Bacteria

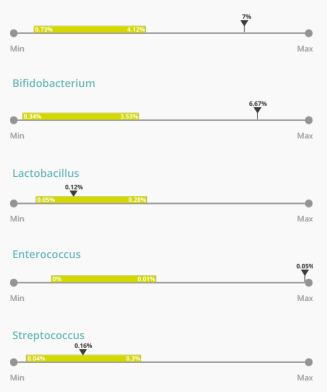


0.02% 0.44% Min Max

Sulphate naturally occurs in our digestion and is, among other things, needed to produce mucins and bile. Sulphate reducing bacteria have specialized in using sulphate for their metabolic processes during which they produce hydrogen sulfide (H2S) A low or average abundance (a few percent) is commonly seen in the microbiome and is not regarded as unhealthy. A high abundance of sulphate reducing bacteria is regarded as unhealthy. Hydrogen sulfide disrupts the cell lining of the intestines (colonocytes) and therefore damages the gut lining. This damage could be the start of an infiammation of the gut and is typically associated with diseases of the gut, such as ulcerative colitis (UC).

Lactate producing bacteria

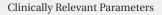
Total number of lactate-forming bacteria



Lactate producing bacteria (also called lactic acid bacteria) are known for their production of lactate. Lactate is an important food source for lactate converting bacteria who cannot live on the sugars from food sources. These bacteria convert lactate into propionate and butyrate (view page 4).

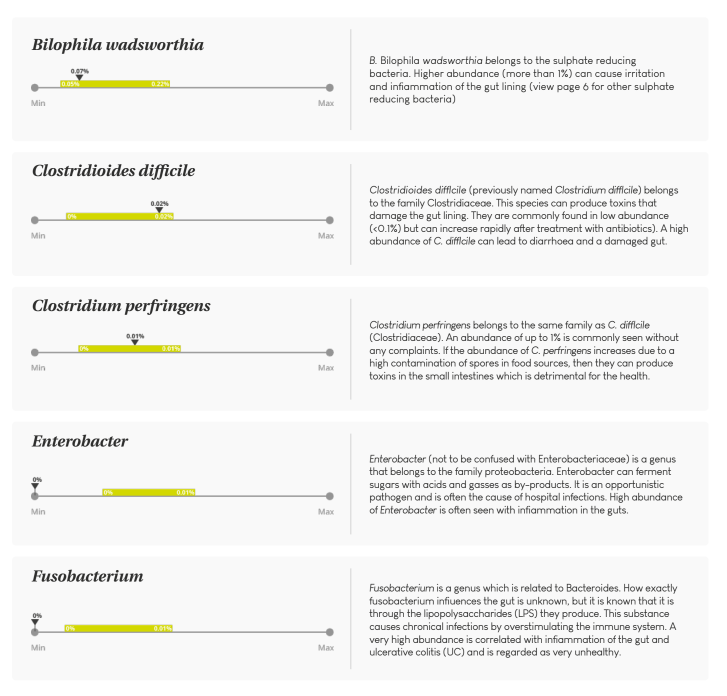
Lactate producing bacteria also produce other metabolites which are known to have a beneficial impact on gut health. These bacteria are often used, due to their ability to produce lactate and other beneficial metabolites, as probiotics. The most well-known groups of lactate producing bacteria are *Bifldobacterium and the Lactobacillus*. However, also *Enterococcus and Streptococcus* are known to produce lactate. A low abundance of lactate producing bacteria can inhibit the production of propionate and butyrate and is therefore regarded as unhealthy (view page 4).

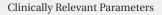
A high abundance of lactate producing bacteria can, when combined with not many butyrate and propionate producing bacteria, result in an excess of lactate in the gut. This can cause several complaints and is mainly seen in people with Crohn's disease or colitis.





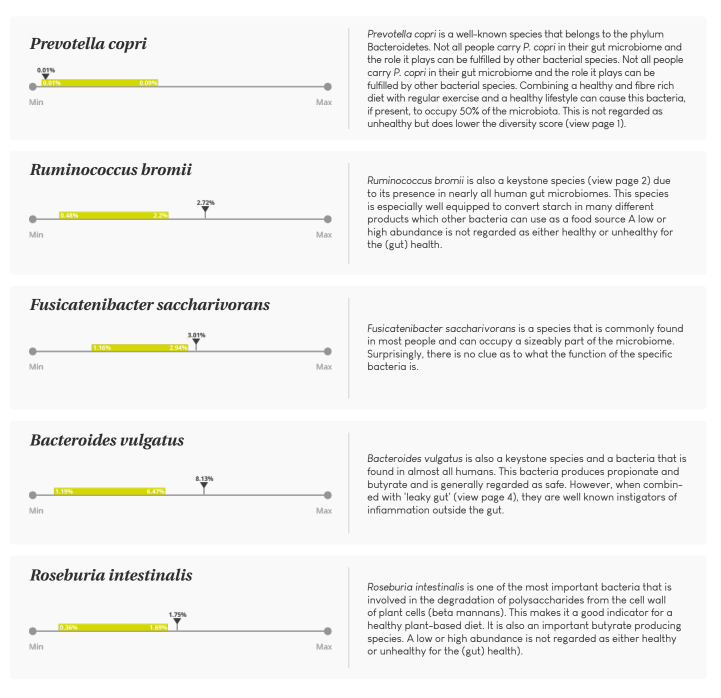
Unwanted species and genera







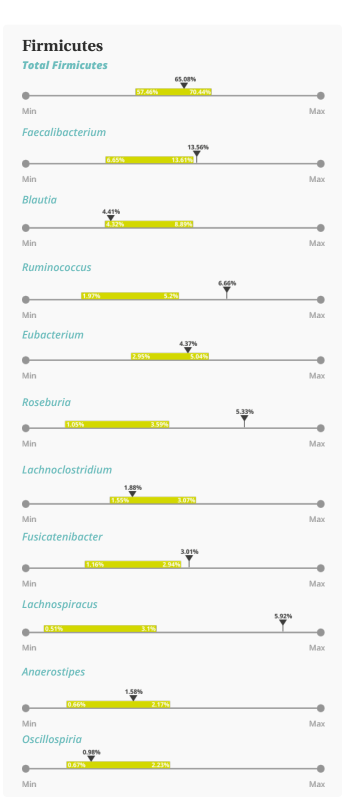
A selection of commonly found species





Genus level ordered by phylum

Note: A selection has been made of relevant genera. Please view the online results to investigate the different taxonomic levels, such as families, genera or species, in more detail.



Clostridium 1.16% . Min Max Phascolarctobacterium • Min Max Megamonas 0% -0 Min Max Coprococcus 0.61% . -0 Min Max Oscillibacter 0.52% a . Min Max Dorea 1.02% • • Min Max Dialister 0.64% ¢ . Min Max Clostridiales 0.03% . -Min Max Christensenella 0% -0 Min Max Acidaminococcus 0.36% • 8 Min Max Lactobacillus 0.12% • -0 Min Max

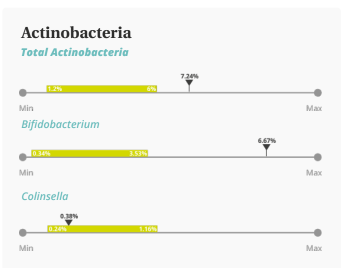
Firmicutes forms the biggest group of bacteria. Most butyrate producing bacteria belong to this phylum.



Bacteroidetes



Bacteroidetes are often the second biggest group of bacteria found in the guts. This group contains many keystone species, which are important for a healthy balance of the gut microbiome.



Actinobacteria are mostly represented by the Bifldobacteria. This genus is a well-known probiotic and one of the most common genus found during in babies and young children. These bacteria thrive on the so-called 'human milk oligosaccharides' (HMO) which are found in breast milk and are important for the development of the immune system This bacteria can also be found in a later stage of life, but it will then function as a lactate producing bacteria (view page 6).



Verrucomicrobia	The only representative of this phyla is the species Akkermansia muciniphila (view page 2). This is one of the keystone species and contributes to a healthy gut and gut microbiome.
Cyanobacteria	Cyanobacteria only account for a small percentage of the gut microbiome. The representatives of this phyla must play a role in the gut microbiome, but we do not know what role yet. It is a possibility that these bacteria live off products produced by other bacteria in small amount or that they thrive on food sources that are scarcely represented in the diet.
Euryarchaeota	<i>Euryarchaeota</i> form an important and special group since these are not bacteria but archaea. Archaea is a kingdom that exists parallel to bacteria. Methanobacteria, who produce methane, are the most well-known representatives of this group.
Proteobacteria	The most important representative of this group is Escherichia coli. This group is also known as an opportunistic pathogen, which are bacteria who only cause illness when new strains are encountered. This is often the cause of food poisoning since new strains are often encountered while visiting new places. A high abundance of prote- obacteria is often not a good sign and is also often found in people with obesity.



Top 10 species

Personal top 10

Faecalibacterium prausnitzii	
6.64% 13.61%	
Min	Max
Bacteroides vulgatus	
8.13%	
1.19% 6.47%	
Min	Max
Eubacterium rectale	
0.45% 3.04%	5.9%
0.45% 3.04%	
Min	Max
Bifidobacterium adolescentis	5.11%
0.12% 1.87%	
Min	
	Max
Bacteroides massiliensis	3.45%
0.01% 1.48%	
Min	Max
Romboutsia sedimentorum	
	3.35%
0.11% 0.75%	
Min	Max
Fusicatenibacter saccharivorans	
3.01% 1.16% 2.94%	
•	
Min	Max
Bacteroides uniformis	
1.07% 3.83%	
Min	Max
Bacteroides stercoris	IVIDA
Dacieroides siercons	2.83%
0.03% 0.59%	
Min	Max
Ruminococcus bromii	
2.72%	
0.48% 2.2%	•
Min	Max