

Article

Exploring Bacterial Communities in Aquaponic Systems

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Abstract: Aquaponics is a production system based on the dynamic equilibrium between fish, plants, and microorganisms. In order to better understand the role of microorganisms in this tripartite relationship, we studied the bacterial communities hosted in eight aquaponic and aquaculture systems. The bacterial communities were analyzed by 16S rRNA gene deep sequencing. At the phylum level, the bacterial communities from all systems were relatively similar with a predominance of *Proteobacteria* and *Bacteroidetes*. At the genus level, however, the communities present in the sampled systems were more heterogeneous. The biofilter samples harbored more diverse communities than the corresponding sump samples. The core microbiomes from the coupled and decoupled systems shared more common operational taxonomic units than with the aquaculture systems. Eventually, some of the taxa identified in the systems could have beneficial functions for plant growth and health, but a deeper analysis would be required to identify the precise functions involved in aquaponics.

Keywords: aquaponics; community analysis; next-generation sequencing; 16S rRNA gene

1. Introduction

Aquaponics is a combination of hydroponic and recirculating aquaculture technologies [1,2]. It offers the possibility of recycling nutrient-rich waste water from fish into organic fertilizers for the plants grown in the system [3], thus reducing the use of fertilizers of mineral origin and the environmental impact of both fish and plant production [4,5]. The use of the aquaculture wastewater to fertilize the plants can avoid the discharge of phosphorus- and nitrogen-enriched water into already nitrogen-loaded surface- and groundwater [5,6].

Along with plants and fish, microorganisms are present in aquaponics. Bacteria are key players in processes which are central for the functioning and equilibrium of an aquaponic system [7]. The best studied process is nitrification, during which ammonia (the main nitrogen form excreted by the fish) is transformed via nitrite to nitrate, which is less toxic for the fish [8] and preferred by plants [9,10]. The main bacteria involved in this transformation are the ammonia oxidizing bacteria (AOB), such as *Nitrosococcus*, *Nitrosospira*, and *Nitrosomonas*, and the nitrite oxidizing bacteria (NOB), such as *Nitrobacter*, *Nitrospira* [11], *Nitrococcus*, and *Nitrospina* [12]. Some *Nitrospira* populations are also able to perform the complete ammonia to nitrate transformation—and are known as complete

ammonia oxidizers (COMAMMOX)—by themselves [13,14]. Archaea, such as the *Thaumarchaeota*, can also be involved in the ammonia-oxidizing process [13]. Finally, the anaerobic ammonium oxidation (ANAMMOX) group, members of the *Planctomycetes* responsible for the anaerobic transformation of ammonium and nitrite into nitrous oxide and N₂ [15] may play a role as well where oxygen levels are low.

In addition to nitrification, microorganisms are involved in other important processes. They can contribute to extracting the various macro- and micronutrients from the feed leftovers and solid feces and make them available for plant uptake [16]. Depending on the aquaculture compartment, design, fish species, and feed type, 15–60% of the consumed feed is actually converted into fish biomass and used for fish metabolism. The rest is excreted and is available for the bacteria to decompose [9,17,18]. Besides this, bacteria could also play a role in the solubilization of nutrients encompassed in solid compounds, such as phytates [19]. Additionally, microorganisms in aquaponics are also involved in various plant growth promotion and protection pathways, such as biocontrol or the enhancement of root growth [7,20,21]. However, these pathways are not sufficiently elucidated yet.

Here, we compared a set of aquaculture (AQ) and aquaponic (AP) systems, which differ in terms of plant and fish species and/or feed type. AP designs included both “coupled” or closed loop AP systems (one loop containing fish and plants) and “decoupled” or open loop AP systems (two separate loops for fish and plants). The aim of this study was to gain insight into the diversity of the bacterial communities in these systems and, if possible, to link their potential functions to plant growth and plant health. For this, the bacterial communities present in biofilter and sump samples, which were the two common units in all systems, were characterized using 16S rRNA gene deep sequencing.

2. Materials and Methods

2.1. Samples Collected in This Study and Samples Preparation

Sump and biofilter samples were collected from eight different systems (Table 1). Three were operated as aquaculture and five as aquaponics. Samples were collected in the period between March and April 2017 as described below. A detailed description of the systems is given in Supplementary Material (Description of the visited aquaponic and aquaculture systems, Figure S1 and Table S1). For comparison to previously published data [7], the dataset of the aquaponic system of the Zürich University of Applied Sciences (ZHAW) was downloaded from the European Bioinformatics Institute database (EBI) and analyzed similarly to the data generated in this study.

2.1.1. Sump Samples

For each sample, two liters of water were collected in sterile Pyrex bottles. In order to concentrate the bacteria, the samples were filtered through 0.2 µm filters (Supor[®])—with a vacuum pump. The filters were then placed in a 50 mL sterile Falcon tube containing 30 mL of sterile water. After vortexing the Falcon tube for 4 min, the filters were removed and the tube centrifuged at 7607 × *g* for 10 min [22]. The pellet was then directly used for DNA extraction.

2.1.2. Biofilter Samples

The biochips used in the different systems varied in shape and size. Therefore, a constant number of biochips per sample could not be taken, as this would not always fit in a 50 mL Falcon tube. Instead, as many biochips as possible (between 10 and 30 depending on the size) were inserted in a 50 mL sterile Falcon tube containing 30 mL of sterile water in order to ensure the harvest of a maximum quantity of bacteria. The Falcon tubes were placed on a vortex for 2 min before placing them for 5 min in an ultrasonic bath (Ultrasonic cleaner, model USC600T, VWR, Leuven, Belgium). The Falcon tubes were then centrifuged at 7607 × *g* for 10 min and the pellet was collected for DNA extraction.

Table 1. Comparison of the sampled aquaculture and aquaponics systems.

Code	Operator	Location	Design	Fish Species	Feed Type	Biochips Type	Plant Type	Sampling Date	Sump Samples	Biofilter Samples	Extra Samples
PCG	Provincial Trial Centre for Vegetable Production	Kruishoutem, Belgium	Aquaponics—open loop	<i>Scrotum barco</i>	Vegetarian	Eco Pondchips	Tomatoes	29/03/2017	1 sump 60 (low density) 1 sump 100 (high density)	1 biofilter 60 (low density) 1 biofilter 100 (high density)	
INA	Inagro	Rumbeke-Beitem, Belgium	Aquaponics—open loop	<i>Sander lucioperca</i>	Omnivorous	Kaldnes media	Tomatoes	18/04/2017	1 sump fish 1 sump hydroponics	1 biofilter	
UF	UrbanFarmers	The Hague, Netherlands	Aquaponics—open loop	<i>Oreochromis niloticus</i>	Omnivorous	Kaldnes media	Microgreens Leafy greens Fruity vegetables	23/03/2017	1 sump	1 biofilter	1 biofilm
IGB	Leibnitz-Institute of freshwater ecology and inland fisheries	Berlin, Germany	Aquaponics—open loop	<i>Oreochromis niloticus</i>	Omnivorous	Kaldnes media	Tomatoes	07/04/2017	1 sump	1 biofilter	
GBXP	Gembloux Agro Bio Tech, PAFF Box system	Gembloux, Belgium	Aquaponics—closed loop	<i>Oreochromis niloticus</i>	Vegetarian	Microbeads	Leafy greens Fruity vegetables	27/04/2017	4 sump	4 biofilter	
GBXR	Gembloux Agro Bio Tech, RAS system	Gembloux, Belgium	Aquaculture	<i>Oreochromis niloticus</i>	Vegetarian	Biocerapond	N.R.	03/04/2017	1 sump	1 biofilter	
BQF	Belgian Quality Fish	Dottignies, Belgium	Aquaculture	<i>Acipenser</i> spp. <i>Huso</i> sp.	Omnivorous	Kaldnes media	N.R.	29/03/2017	1 sump	1 biofilter	1 biofilter denitrification
WU	Wageningen University	Wageningen, Netherlands	Aquaculture	Eel, catfish	Omnivorous	Kaldnes media	N.R.	12/04/2017	1 sump eel 1 sump catfish	1 biofilter eel	

N.R.: not relevant.

2.2. DNA Extraction

The Fast DNA Spin Kit using Cell Lysis Solution TC (MP Biomedicals, Santa Ana, CA, USA) was used for the DNA extractions. The manufacturer's protocol was modified as follows: Samples were homogenised with a Power-Mix Model L46 (Labinco, Breda, The Netherlands) at speed setting 7 for 40 s, then incubated on ice for 2 min and again homogenized for 40 s. Subsequently, to remove cell debris, tubes were centrifuged at $14,000\times g$ for 10 min. All DNA extracts were stored unopened at $4\text{ }^{\circ}\text{C}$ until further analyses.

2.3. Sequencing

Library preparation and sequencing were carried out by DNA Vision S.A. (company, Gosselies, Belgium). The library preparation and indexing steps were done using the Nextera Index kit v2 (Illumina, San Diego, CA, USA), while sequencing was conducted on an Illumina Miseq (2×250 bp) (Illumina Inc., San Diego, CA, USA) with the Miseq reagent kit v3 (600-cycles, Illumina, San Diego, CA, USA). Sequencing primers were chosen to cover the hypervariable regions V1-V3 of the 16S ribosomal RNA gene as recommended by Munguia-Fragozo et al. [23] and Schmutz et al. [7]. The following primers were used:

Forward V1-V3

5'-TCGTCGGCAGCGTCAGATGTGTATAAGAGACAGAGAGTTTGATCCTGGCTCAG-3'

Reverse V1-V3

5'-GTCTCGTGGGCTCGGAGATGTGTATAAGAGACAGATTACCGCGGCTGCTGG-3'

Data are available under the accession PRJNA513832 on the Sequence Read Archive database (SRA) of the National Center for Biotechnology Information (NCBI).

2.4. Bioinformatics

The analysis of the sequencing data was conducted with the QIIME pipeline v1.9.1 (<http://qiime.org/>) [24]. Forward and reverse sequences were merged in one file per sample with `multiple_join_paired_ends.py`. Paired fastq files were converted into fasta files with `convert_fastaqual_fastq.py`. Sequences were assigned to Operational Taxonomic Units (OTUs) with a cut-off of 97% sequence identity to the reference database Greengenes 13_8 with `pick_de_novo_otus.py`. Chimeric sequences were identified with the Chimera Slayer tool and then removed with `filter_fasta.py`. Singletons and sequences originating from chloroplasts and mitochondria were discarded.

For further analysis of the bacteria communities' composition in the 22 samples, the samples were rarefied at 40,000 sequences with `single_rarefaction.py` (rarefaction curves available in the Supplementary Material; Figure S2). Bar charts representing the relative abundances of the various OTUs were obtained using `summarize_taxa_through_plots.py`. Core microbiomes were generated using `compute_core_microbiome.py`.

2.5. Statistics

The Shannon and equitability indices were calculated via the `alpha_diversity.py`. The Shannon and equitability indices of biofilter versus sump were compared via the `global_core_diversity_analyses.py` workflow with a nonparametric t-test (using Monte Carlo permutations). Principal coordinates analyses were carried out with the `beta_diversity_through_plots.py` script in order to compare the communities.

3. Results

3.1. Metagenome Sequencing

Whereas the previous study on the Wädenswil Aquaponics System [7] gave a first impression on the bacterial communities in a single setting, a comparison to other systems was not performed. For this

reason, we collected 22 samples from various aquaponic and aquaculture systems in Western Europe (Table 1). Eleven samples were collected from sumps, nine from biofilters, one from an additional denitrification biofilter, and one from the periphyton present on tank walls (Table 2). The bacterial communities thereof were analyzed in this study using 16S rRNA gene deep sequencing. The average Q_{30} of the sequence in the samples was 80%, which indicated samples of good DNA quality. Of the total reads, 11.8% were not assigned at the phylum level.

3.2. Taxonomic Assignment of Reads

Based on the taxonomic assignment of the reads, it can be observed that the bacterial communities in the different systems were highly variable. Of all filtered reads in all samples, an average $11.8\% \pm 6.7\%$ could not be assigned to any OTU. In general, two major phyla were found throughout the samples (Figure 1): *Proteobacteria*, representing $34.6\% \pm 10.1\%$ of the total reads, and *Bacteroidetes*, representing $25.5\% \pm 14.0\%$. Other phyla were found in lower quantities in the samples. However, some samples held exceptionally high amounts of individual phyla. An example here is the presence of 73.1% reads representing the *Thermi* phylum, mainly represented by a single OTU (*Deinococcus*) in the sump sample of the Belgian Quality Fish (BQF) system (Figure 1). Except for the biofilter of the same system, this phylum was only present at very low levels in the other systems.

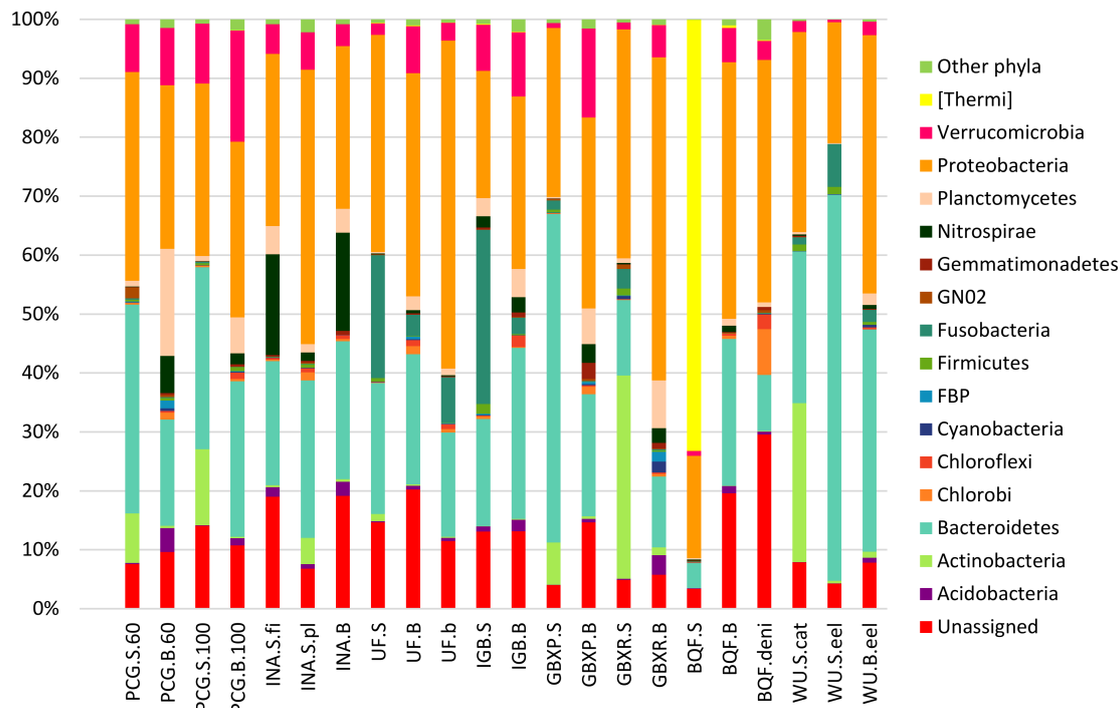


Figure 1. Bar charts representing the relative abundances of the phyla in each sample. Phyla which represented less than 0.2% of the total reads are gathered under “other phyla” *BHI 80139*, *BRC1*, *Chlamydia*, *Elusimicrobia*, *Fibrobacteras*, *GN04*, *GOUTA4*, *Lentispaerae*, *NKB19*, *OP11*, *OP3*, *OP8*, *PAUC34f*, *SBR1093*, *SR1*, *Spirochaetes*, *Synergistetes*, *TM6*, *TM7*, *Tenericutes*, *WPS2*, *WS1*, *WS2*, *WS3*, *WWE1*, and *Caldithrix*).

Table 2. Summary of metagenomics data.

Code	Sampling Zone	Number of Reads before Filtering	Chimeric Reads	Chloroplast and Mitochondrial Reads	Singleton Reads	Number of Reads after Filtering	% of Unassigned Reads	Shannon Index (after Filtering)	Equitability Index (after Filtering)
PCG.S.60	sump low density	75,840	83	9	1016	74,732	7.6%	5.65	0.52
PCG.S.100	sump high density	131,231	443	100	3822	127,166	13.9%	6.55	0.56
PCG.B.60	biofilter low density	104,241	77	10	2661	101,493	9.5%	7.88	0.69
PCG.B.100	biofilter high density	78,392	66	7	1635	76,684	10.6%	7.73	0.70
INA.S.fi	sump fish loop	107,998	146	117	2634	105,101	19.0%	7.68	0.66
INA.S.pl	sump before plant compartment	92,790	708	2	3581	87,985	6.8%	8.63	0.71
INA.B	biofilter	100,948	154	124	2831	97,839	19.2%	7.37	0.65
UES	sump	117,695	1122	22	3175	113,376	14.6%	6.77	0.57
UEB	biofilter	134,730	223	5	7336	126,866	20.3%	8.30	0.67
UEb	biofilm	99,487	260	56	2407	96,764	11.4%	6.94	0.59
IGB.S	sump	63,482	186	10	2657	57,612	12.9%	7.50	0.62
IGB.B	biofilter	59,923	74	45	2192	58,091	13.1%	8.44	0.75
GBXP.S	sump	97,905	340	864	4	96,697	4.0%	3.79	0.34
GBXP.B	biofilter	69,831	52	0	2233	67,546	14.7%	8.31	0.74
GBXR.S	sump	11,096	1037	100	2318	112,641	4.8%	5.91	0.51
GBXR.B	biofilter	124,569	378	81	3937	120,173	5.7%	7.65	0.65
BQES	sump	81,204	50	10	1087	80,057	3.3%	2.83	0.26
BQFB	biofilter	56,448	85	13	3090	53,260	19.7%	8.42	0.74
BQF.deni	denitrification biofilter	65,693	14	1	1966	63,712	29.5%	6.72	0.63
WU.S.cat	sump catfish system	44,743	119	4	1142	43,478	7.9%	6.20	0.58
WU.S.eel	sump eel system	75,055	32	1	671	74,351	4.2%	3.22	0.32
WU.B.eel	biofilter eel system	101,169	177	8	2059	98,925	7.8%	6.43	0.60

It could be noted that in almost all systems, the biofilter sample harbored a more diverse community as the Shannon indices of the biofilter samples were significantly higher ($p < 0.05$) than the Shannon indices of the sump group (Table 2). The equitability was also significantly higher in the biofilter samples than is the sump samples ($p < 0.05$). The only exception was the INA system. A more thorough exploration of this system would be required to explain this difference.

At the genus level, reads were assigned to more than 700 different OTUs. To allow for a more in-depth analysis of the genera present in aquaponic systems, it was decided to focus on the OTUs representing more than 1% of the total reads per sample (Figure 2). For some OTUs, the identification process was only possible at the family level.

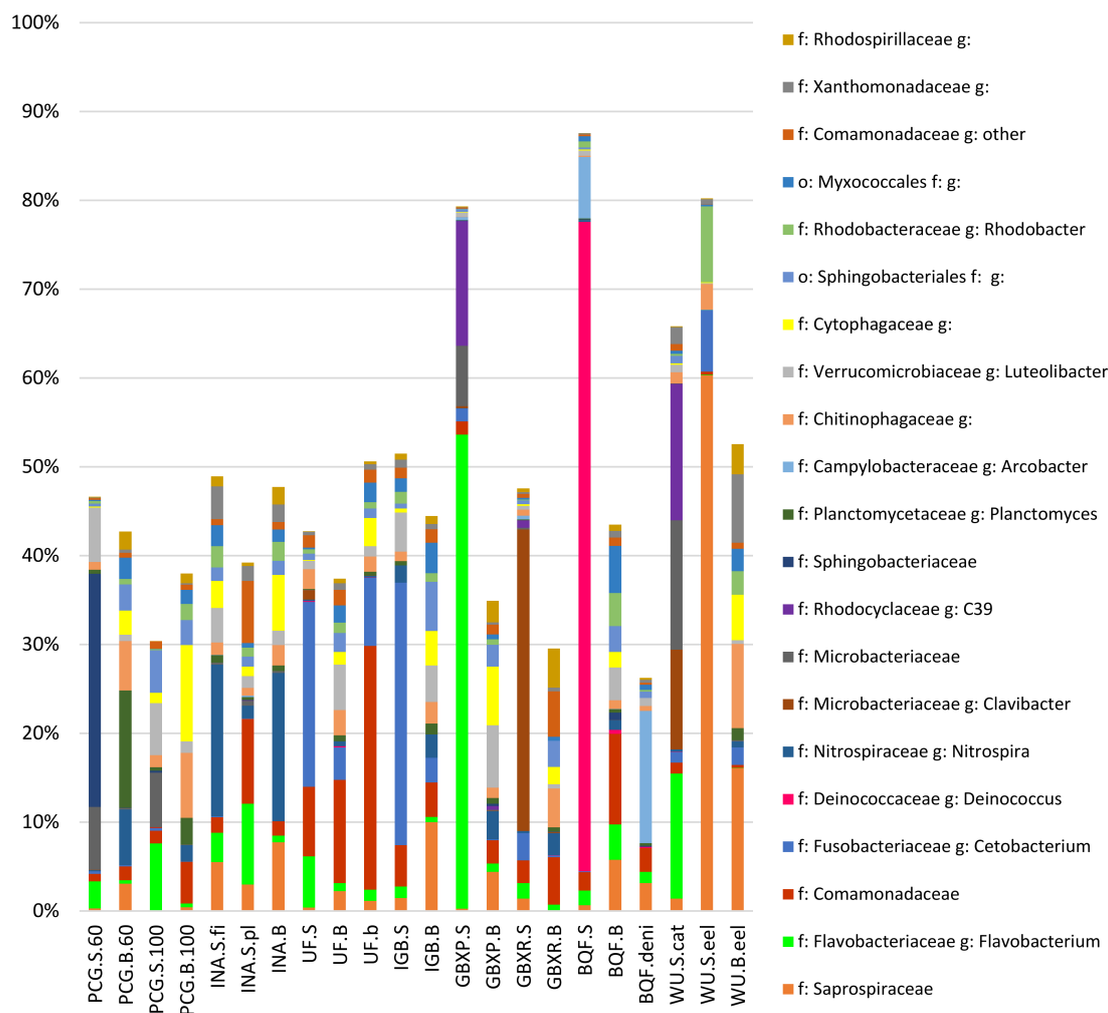


Figure 2. Bar charts representing the relative abundances of the families and genera representing more than 1% of the total reads for each sample.

3.3. Potential Influence of the System Design on Microbial Communities

One OTU assigned to the genus *Deinococcus* was mainly present in the BQF system and represented 73% of the BQF sump sample. Members of the genus *Deinococcus* are heterotrophic organisms resistant to UV radiation [25] and to our knowledge, there is no link between the rearing of sturgeons and the presence of this genus. As the BQF system was implemented with an ozone plus UV light disinfection treatment (Supplementary Material, Description of the visited aquaponic and aquaculture systems), it could be expected that this organism was dominating the community based on its resistance to such treatment. However, most aquaponic systems, such as the Wädenswil Aquaponics system [7], use UV light to prevent the proliferation of undesirable microorganisms and to keep the water clean and

clear [9], without observing such a development of *Deinococcus*. It could thus be that the dominance of *Deinococcus spp.* in the sump was due to a higher strength of the UV light in combination with the ozone treatment. It was also observed in both biofilter samples, but at levels below 0.5% of the community. This indicated that the *Deinococcus spp.* were more planktonic, while we observed a broader diversity in the biofilms on the carrier in the biofilters.

The species *Cetobacterium somerae* belongs to the *Fusobacteriaceae* family and has been commonly found in guts of freshwater fish [7,12,26]. The discovery of *Cetobacterium* in some of the samples might indicate that the system design in these cases was not sufficiently adapted to remove sufficient amounts of fish feces from the water of the fish tank. On the day when the Urban Farmers (UF) (Supplementary Material, Description of the visited aquaponic and aquaculture systems) samples were collected, pipes were clogged with fish sludge in the drum filter compartment and thus sludge may have passed the drum filter towards the biofilter and sump compartments. This might explain the large amounts of reads assigned to the genus *Cetobacterium* in the UF samples (Figure 2). Additionally, it was observed that *Cetobacterium* were more often detected in the sump of systems than in biofilter samples. It was assumed that the sump in these systems could offer sufficiently anaerobic zones whereas the moving bed biofilters were fully aerobic [3]. However, if the system hosted an important quantity of *Cetobacterium* in the sump, one could also observe their presence in the biofilter albeit at a smaller proportion (Figure 2).

Conversion of nitrogen compounds is of utmost importance for recirculating systems to avoid toxicity problems of the different nitrogen forms for each species. Of the known nitrifying bacteria, the *Nitrosomonadaceae* family was present in most samples. Even though the relative abundance of this family was quite low (between 0% and 1.7% of reads; average = 0.3%; stdev = 0.5%), the order of magnitude observed in most samples in this study was similar to the one observed in the study of the Wädenswil Aquaponics system [7]. The most abundant nitrifying bacteria were those of the genus *Nitrospira*. This is also in accordance with the study conducted on the Wädenswil Aquaponics system [7], and may indicate that the COMAMMOX process is more common to aquatic culture systems.

3.4. Core Microbiomes

3.4.1. General Core Microbiome

Generally, a large diversity of bacteria was observed and all systems hosted different bacterial communities (Figure 1). However, in spite of this diversity and the specificities of each system, common bacterial groups were found in all aquaponic and aquaculture systems. A core microbiome containing only the OTUs present in all samples was extracted from the data set and, regardless of the system and sample location, four OTUs were identified. OTUs representing unidentified genera from the *Oxalobacteraceae* family and the *Comamonadaceae* family were identified as being present in all samples. The *Oxalobacteraceae* family harbors several heterotrophic bacteria that can be found in water, soil and also in association with plants [27]. Regarding the *Comamonadaceae* family, it is also found in aquaculture or aquaponic systems in other studies [7,12]. In their review, Munguia-Fragoso et al. [23] reported that *Comamonas sp.* were identified in several bacterial communities of freshwater recirculating aquaculture systems (RAS). At the genus level, OTUs assigned to the genera *Cetobacterium* were as well part of the general core microbiome. Although *Cetobacterium* is rather an anaerobe, its presence in all systems could be explained by its common presence in fish guts [26,28].

3.4.2. System-Specific Core Microbiomes

On the basis of the three basic setups that were sampled in this study, we also calculated core microbiomes for each of these setups (Figure 3). The AQ group contained nine common OTUs, and the decoupled AP group harbored a core microbiome of 34 OTUs. The coupled AP group contained only the plant and fish farming (PAFF) Box samples and, therefore, harbored a core microbiome of 636 OTUs. Whilst only one and five OTUs were common for aquaculture and the two aquaponics

systems, the two aquaponic groups share more OTUs. A total of 17 OTUs at different levels were found, indicating that there are some not yet identified conditions that are specific to aquaponic systems, independently of the setup.

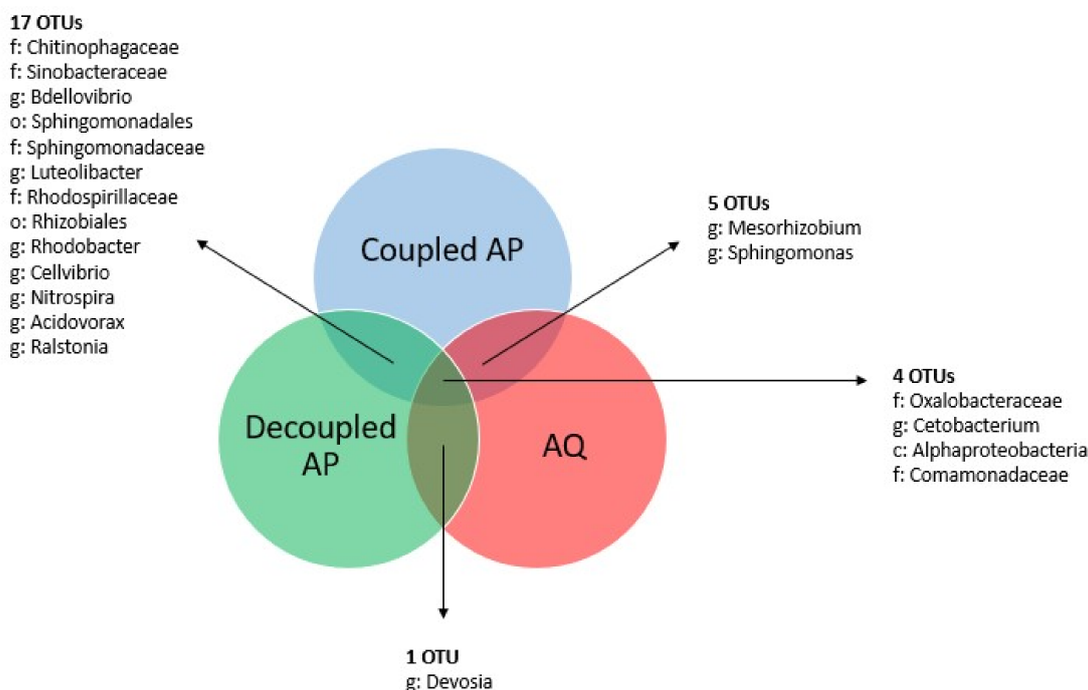


Figure 3. Venn diagram representing the intersection of different core microbiomes obtained through the grouping of samples based on the system setup (i.e., coupled aquaponics, decoupled aquaponics, and aquaculture).

3.4.3. Sampling Site-Specific Core Microbiomes

Based on the sampling strategy, sampling site-specific core microbiomes were generated from biofilter and sump samples. The sump core microbiome was composed of 22 OTUs, while the biofilter core microbiome was composed of 28 OTUs. The larger numbers of OTUs in site-specific core microbiomes indicated that both zones had site-specific bacterial communities. This was confirmed by principal coordinates analysis (PcoA), indicating the presence of two sample clusters: a narrow cluster grouping biofilter samples and a wider cluster grouping the sump samples (Figure 4). Between these two site-specific core microbiomes, ten OTUs were common (four of them belonging to the global core microbiome). The six additional OTUs found in all biofilters and sumps were *Sphingomonas*, *Devosia*, *Novosphingobium*, *Acidovorax*, *Ralstonia*, and an unidentified OTU from the *Rhizobiaceae* family. *Nitrospira* could be found in all biofilter samples. However, this was not the case for the genera *Nitrosomonas* and *Nitrobacter*.

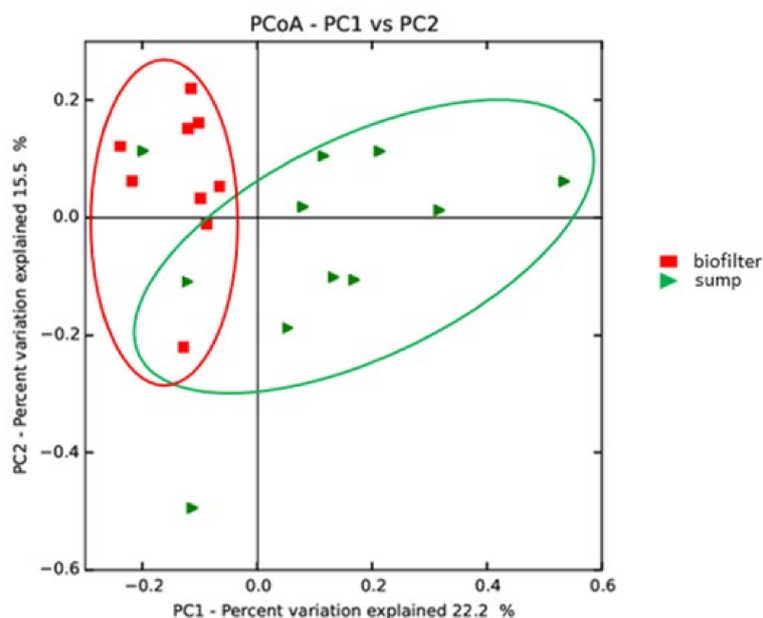


Figure 4. Weighted UniFrac principal coordinates analysis presenting the separation between the sumps and the biofilter samples. Axis 1 and 2 explain 37.8% of the total variability.

3.5. Detailed Analysis of the Microbial Communities in the Coupled and Decoupled Gembloux Systems

Up to now, we have compared systems that were highly heterogeneous in their design, size, and operational strategies. This study also included two systems that were highly comparable: the recirculating aquaculture system (RAS) and the Plant and Fish Farming (PAFF) Box of Gembloux Agro–Bio Tech, as both systems shared the same size, fish, feed type and incoming water quality. The main difference was that the PAFF Box is operated as closed circular system, whereas the RAS is an aquaculture system. In order to observe the impact of plants in the system on the composition of the bacterial community, we chose to compare the communities in these two systems in more detail.

The RAS sump sample was dominated by the two genera *Clavibacter* and *Cetobacterium*, whereas the PAFF Box sample contained a majority of the genus *Flavobacterium* and C39, an OTU belonging to the *Rhodocyclaceae* family (Figure 5). The presence of such large numbers of reads assigned to *Clavibacter*, a genus that mainly contains plant pathogenic species [29], needs to be examined in more detail to confirm the assignment of all reads to this genus. The RAS biofilter sample was clearly dominated by members of the genus *Lysobacter* and also hosted *Nitrospira* and *Novosphingobium*, while the PAFF Box biofilter contained C39, *Nitrospira*, *Flavobacterium*, members of the *Microbacteriaceae* family, and *Cetobacterium* (Figure 5). This indicated that, despite similarities in the global setup (fish species, feed type, incoming water quality, and size of the fish tanks), each system developed its own specific community. The presence of plants in the aquaculture loop thus had a large influence on the composition of the bacterial community.

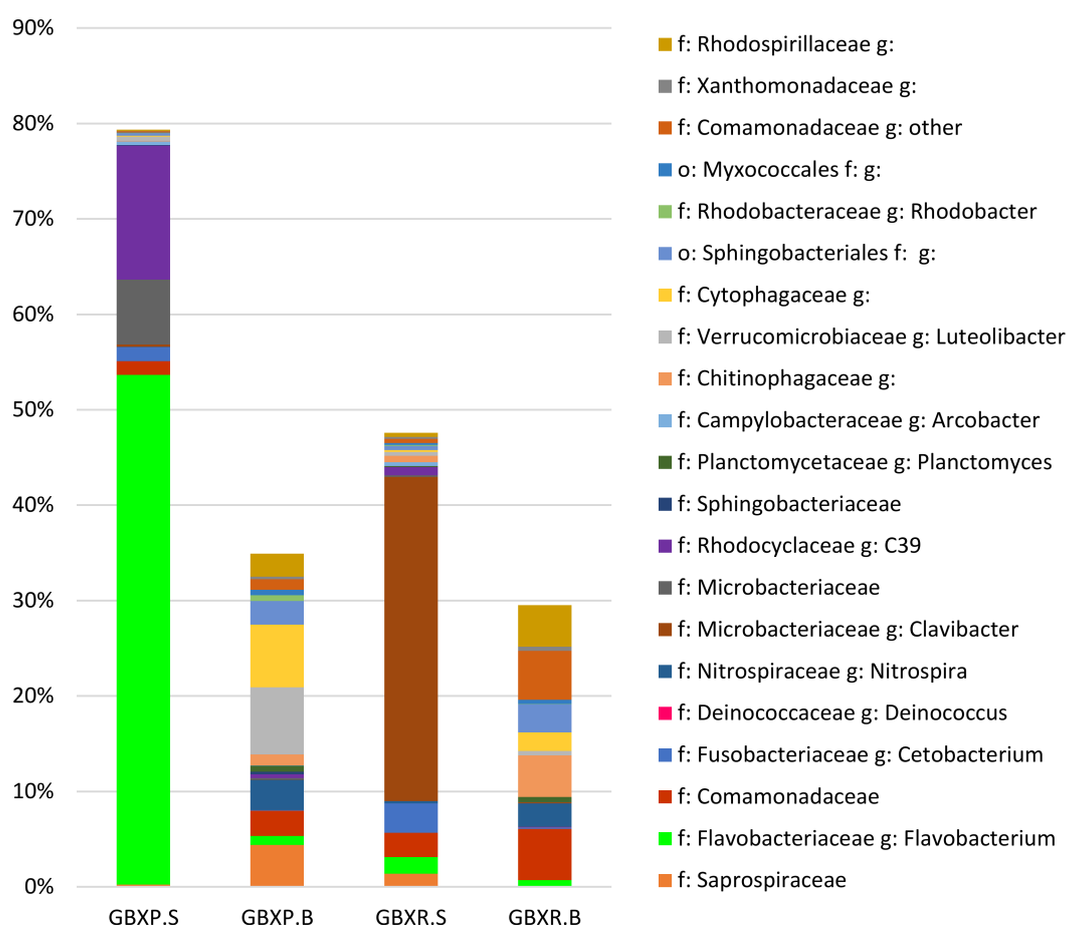


Figure 5. Relative abundances of the families and genera representing more than 1% of the total reads within each sample, for the biofilter and sump samples of the recirculating aquaculture system (RAS) and plant and fish farming (PAFF) Box systems of Gembloux Agro-Bio. Tech.

4. Discussion

4.1. Predominant Taxa

In order to work towards better understanding of how bacterial communities function in aquaponics, we decided to focus on the potential impact of the bacteria in the aquaponic solution on plant growth. A few studies have reported that, with aquaponics, they obtained plant yields as good as in hydroponics despite the aquaponic solution containing lower nutrient concentrations [1,8,30]. A first step to take towards the elucidation of this increased growth would be to check for similarities in the bacterial communities in different aquaponic systems. In order to do so, it was decided to study the core microbiomes of our samples. The concept of core microbiomes has been used and described in several other research fields, such as the plant holobiont [31], humans [32], or milk microbiomes [33]. The core microbiome has been defined by Lemanceau et al. [31] (p. 1) as “the microbial community that is systematically associated with a given host”. Until now, the concept of core microbiome has been focused on the taxonomic composition of a community. However, it could also be argued that a core microbiome should have specific functionalities responding to the needs of their associated host [31].

At the phylum level, *Proteobacteria* and *Bacteroidetes* were the two major groups representing, respectively, 35% and 26% of the reads obtained from the different aquaponics and aquaculture systems. This was in accordance with the observation from Schmutz et al. [7], who also found that *Proteobacteria* (approximately 50%) and *Bacteroidetes* (15–20%) were the major phyla in their root

zone, biofilter, and periphyton samples. This also corroborated observations made in freshwater aquaculture [23]. Other phyla common to the observations by Schmautz et al. [7], the freshwater data cited by Munguia-Fragoso et al. [23], and the present study were the phyla *Actinobacteria*, *Planctomycetes* and *Nitrospirae*.

Few reads assigned to the genus *Pseudomonas* were detected in the visited systems. The number of assigned reads to the pseudomonads was in the same order of magnitude as previously observed by Schmautz et al. [7]. *Pseudomonas* are usually found in close proximity to roots and are less prevalent in bulk soil [34]. As, in this study, we did not investigate the root zone of the aquaponics systems, this genus may thus rather be represented by planktonic species in the samples taken.

4.2. Potential Roles/Functions of the Identified Taxa

In aquaponics, we are interested in the bacteria, which could help us ensure fish welfare and plant care. When it comes to plants, bacteria could help their growth and health through growth stimulation and biocontrol. Taxa, such as the *Microbacteriaceae* family, are known to be able to form associations with plants [35] and have been detected in the microbiome of barley roots along with members of the *Comamonadaceae* family [36]. In the latter family, several species have been detected to possess skills for siderophore production and protection against *Fusarium* and *Rhizoctonia* [37]. The *Microbacteriaceae* family contains species which have aminocyclopropane-1-carboxylate (ACC) deaminase activity, siderophore, and indole production [37]. The *Flavobacterium* genus is widely present in nature and mostly known for its capacity to degrade complex organic molecules [38,39]. *Flavobacterium spp.* are often found in association with plant roots and plant leaves and are believed to be involved in plant growth and protection [38–40]. Several strains were detected to be able to participate in the solubilization of insoluble phosphate, the production of auxin, and the production of siderophore [37,40]. *Flavobacterium* are also used to fight against plant pathogens (*Phytophthora infestans*) in biological control formulation [41]. In aquaculture and aquaponics, *Flavobacterium* have also been detected and can be considered as a common genus found in such systems [7,12,23]. The *Lysobacter* genus has also been identified as plant growth promoting bacteria (PGPB) and can help fight against plant disease through the production of antibiotics [42–44]. The presence of all of those species in most of the samples in this study confirmed the data from the samples from the Wädenswill aquaponics system [7] and strengthens the statement that the microbiome in aquaponics or aquaculture systems may be able to secure the health and growth of the plants in the first-named type of systems.

A crucial function of the bacteria communities in aquaponics would be the solubilization of the fish dejections and fish feed leftovers into macro- and micronutrients, which the plants can absorb. The members of the genera *Flavobacterium* and *Sphingobacterium* could participate in the decomposition of organic matter [39]. The *Saprospiraceae* family is typically found in aquatic environment, such as wastewater treatment plants [39,45,46], and could be involved in the degradation of complex carbon molecules, such as proteins [45,46]. Many other detected genera include heterotrophic organisms able to degrade biomass in the system, as well.

Several of the observed OTUs were related to a role in the nitrogen cycle. The genus *Nitrospira* was detected in all biofilter samples. *Nitrospira* is commonly known as a NOB [11,12,14,47]. Daims et al. [14] showed that certain strains of the *Nitrospira* genus could actually be complete nitrifiers, i.e., able to oxidize ammonia to nitrate without the help of AOB, a process now known under the name COMAMMOX. Denitrification has also been often observed in aquaculture and aquaponics [48,49]. Members of the genus *Arcobacter* are known to perform denitrification [50] and have been particularly found in the denitrifying biofilter of the BQF system (Figure 2). The phylum *Planctomycetes* has already been observed in recirculating aquaculture [51] and contains ANAMMOX bacteria [39,51]. However, with the current database used for assignment to genus level, it could not be confirmed that ANAMMOX bacteria were present in the examined systems.

4.3. Core Microbiomes

Bacterial populations in aquaponic systems were highly diverse whether between systems or between the different compartments therein. In each sample, all the taxa representing less than 1% of the total number of reads were discarded, and this represented at least 50% of the reads (Figure 2). Despite this diversity, a core microbiome common to all samples could be identified. Moreover, the core microbiomes were composed of 28 (7.6% of the total reads common to all biofilters) and 22 OTUs (6.1% of the total reads are common to all sumps) for the biofilter and sump samples, respectively. This brought forth the hypothesis that a common bacterial base may exist between all aquaponic systems despite differences in fish species, system layout, or fish feed.

Despite the differences in the bacterial communities due to system specificities, there were still similarities between all examined systems. A principal coordinate analysis combining the data from the tested systems with the data collected by Schmautz et al. [7] (Supplementary Material; Figure S3) showed that the samples collected from the plant roots, periphyton, and biofilter compartments of the Wädenswil Aquaponic system (ZHAW) clustered closely with the other samples. This showed that there was a common pattern concerning the composition of the bacteria community in diversified aquaponic systems located in Western Europe. It would then be interesting to broaden the study to systems situated worldwide and also on a longer period of time.

5. Conclusion and Perspectives

This study was one of the first investigations into the diversity of bacterial communities present in a variety of aquaponic and aquaculture systems. It offered a global overview of the microbial taxa therein and of the potential roles that microorganisms could play in plant care. Nevertheless, it was shown that the different system setups had a large influence on the bacterial communities, and it needs to be investigated in more detail which species performs what role in such systems.

As the currently available datasets were from a single time point and only limited compartments within single systems were sampled, a more comprehensive sampling of single systems over time would be required to study the influence of sample time and location within a system. This may explain the currently obtained data better within the frame of the operational differences, but helps us also to understand the biological processes taking place in a single system.

Supplementary Materials: The following are available online at <http://www.mdpi.com/2073-4441/11/2/260/s1>, Supplemental material containing a description of the visited aquaponic and aquaculture systems, Table S1: Water quality parameters, Figure S1: Recirculating aquaculture system of the Integrated and Urban Plant Pathology Laboratory, Figure S2: Rarefaction curves of every samples indicating the number of observed OTUs, according to sequencing depth, Figure S3: Weighted UniFrac Principal Coordinates Analysis, including the eight visited systems and the Wädenswill Aquaponic System.

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