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# Inter Individual Variations of the Fish Skin Microbiota: Host Genetics Basis of Mutualism?



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# Abstract

The commensal microbiota of fish skin is suspected to provide a protection against opportunist infections. The skin of fish harbors a complex and diverse microbiota that closely interacts with the surrounding water microbial communities. Up to now there is no clear evidence as to whether the host regulates the recruitment of environmental bacteria to build a specific skin microbiota. To address this question, we detected Quantitative Trait Loci (QTL) associated with the abundance of specific skin microbiota bacterial strains in brook charr (*Salvelinus fontinalis*), combining 16S RNA tagged-amplicon 454 pyrosequencing with genetic linkage analysis. Skin microbiota analysis revealed high inter-individual variation among 86 F2 fish progeny based upon the relative abundance of bacterial operational taxonomic units (OTUs). Out of those OTUs, the pathogenic strain *Flavobacterium psychrophilum* and the non-pathogenic strain *Methylobacterium rhodesianum* explained the majority of inter-individual distances. Furthermore, a strong negative correlation was found between *Flavobacterium* and *Methylobacterium*, suggesting a mutually competitive relationship. Finally, after considering a total of 266 markers, genetic linkage analysis highlighted three major QTL associated with the abundance of *Lysobacter, Rheinheimera* and *Methylobacterium*. All these three genera are known for their beneficial antibacterial activity. Overall, our results provide evidence that host genotype may regulate the abundance of specific genera among their surface microbiota. They also indicate that *Lysobacter, Rheinheimera* and *Methylobacterium* are potentially important genera in providing protection against pathogens.

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## Introduction

The study of the beneficial effects of bacteria and their influence on host health is a growing field. Namely, research that has explored host microbiota variability in space and time suggests the presence of a host genetic component into the development of mutualism communities [1,2]. Similar conclusions were found in several studies on twins and on their core gut microbiota [3,4,5,6]. Furthermore, microbiome homeostasis seems to be the key to resistance against some diseases previously considered exclusively influenced by genetic factors [7]. To determine precisely which genes are involved in the recruitment of specific bacterial strains, some studies looked at gene expression in presence of symbiotic bacteria. Recent studies specifically targeted genes already known as being involved in innate or adaptive immunity (for example IgA) [8,9,10].

In fish, the influence of host genetic background on commensal bacterial community structure is poorly known. However, advances in the field of probiotic development indicate that endogenous bacteria are able to outcompete pathogens [11,12]. Results obtained in aquaculture settings are consistent with previous work showing that non-pathogenic bacteria associated

with animal mucosa can contribute to the host health by providing protection against pathogen infections [13,14,15,16]. The genetic basis of the host immune response, especially at the major histocompatibility complex (MHC), toll-like receptor (TLR) and immunoglobulin loci, has been well studied [17,18,19,20]. However, aside from the immune response, the influence of the host on the development of bacterial symbiosis remains poorly understood.

Here, the main objective of the present work was to investigate the potential link between host genotype and skin microbiota composition. As a host model, we focused on brook charr (*Salvelinus fontinalis*), a salmonid that harbors a complex and dynamic skin microbiota [21,22]. The first specific objective sought to document the genetic variation present in the host population that might underpin the inter-individual variations in the skin microbiota. The second specific objective was to characterize the relationship (cooperation/competition) existing among the different bacterial Operational taxonomic units (OTU) within the skin mucus, which influences the overall structure of the skin microbial community. The third objective was to identify host genetic regions associated to the abundance of specific skin bacterial OTUs (i. e. quantitative trait loci (QTL) associated with 
 Table 1. Descriptive statistics of sequencing to determine the microbiotoa for each individual.

Sample ID	number of reads	coverage	npshannon index	Number of OTU total	Number of OTU >10 reads
92	229	0.955975	1.690045	76	19
93	1584	0.99131	0.436879	74	28
94	669	0.977401	1.507996	242	20
95	738	0.986166	1.416086	226	27
96	503	0.953079	1.395581	149	29
97	385	0.946203	1.774329	86	33
98	955	0.98389	0.663968	79	30
99	770	0.978852	1.306709	114	29
100	1018	0.985946	0.658787	107	30
101	1482	0.994792	0.842304	188	21
102	1667	0.992283	0.691189	96	31
103	711	0.99179	0.946539	97	23
104	1072	0.987599	0.719994	159	25
105	1662	0.991645	0.371924	73	26
106	1304	0.995734	0.658557	93	18
107	1617	0.99416	0.290016	49	21
108	1237	0.990451	0.634742	73	30
109	1063	0.987778	0.679124	137	26
110	752	0.97281	0.656197	105	29
111	731	0.986217	0.63276	79	22
112	628	0.975779	0.630137	63	23
113	892	0.983549	0.446389	49	24
114	1333	0.993127	0.713271	127	22
115	1109	0.983313	1.020009	266	35
116	532	0.967611	2.709733	269	32
117	649	0.963145	2.108945	237	41
118	404	0.923977	2.645631	222	29
119	969	0.97625	1.019139	177	40
120	581	0.967647	1.975631	205	26
121	468	0.948413	2.692236	202	34
122	744	0.980551	0.876389	117	27
123	953	0.983204	0.832745	133	32
124	530	0.967846	2.29117	186	30
125	797	0.977901	0.703269	79	30
126	411	0.97561	0.904985	116	18
120	242	0.76	3.452235	182	32
128	1230	0.989637	0.504536	69	26
120	831	0.978723	1.537117	115	41
130	1449	0.983421	0.548428	123	38
	499	0.969697	2.118408	123	29
31	499	0.969697		144	29
32			2.133064		
33	494	0.96114	1.408929	117	35
34	746	0.980254	0.508333	38	22
135	662	0.977887	2.41624	180	36
36	754	0.987755	0.956343	118	18
137	2008	0.993154	0.557917	74	30
138	714	0.976619	1.632525	101	36
39	798 469	0.967632	1.534968	129 261	35 35

#### Table 1. Cont.

Sample ID	number of reads	coverage	npshannon index	Number of OTU total	Number of OTU >10 reads
141	224	0.980892	0.662835	33	6
142	517	0.957377	2.388226	180	36
143	483	0.947195	2.446711	157	36
144	414	0.958848	3.124298	188	38
145	930	0.975443	1.391363	168	43
146	769	0.979661	2.024708	140	28
147	1219	0.995122	1.641489	118	17
148	1254	0.98244	1.127563	148	40
149	1074	0.972678	2.309049	318	55
150	1108	0.980583	2.793443	321	47
151	3082	0.998643	0.332436	58	16
152	1429	0.996561	1.820656	203	25
153	1955	0.995286	0.815557	167	27
154	1449	0.983366	1.74373	354	54
155	1074	0.973611	1.716763	321	41
156	1016	0.989316	0.588719	62	21
157	1483	0.984836	1.374063	216	46
158	2004	0.998944	0.791157	44	10
159	2043	0.997359	0.812388	92	18
160	2468	0.995741	0.305525	70	23
161	1215	0.979566	1.928985	346	48
162	1904	0.995286	0.821609	164	37
163	1046	0.985799	1.569251	188	40
164	1145	0.993814	0.744587	98	19
165	995	0.981481	1.421104	243	37
166	725	0.97921	2.459141	211	36
167	727	0.988743	2.142562	131	24
168	841	0.98908	2.297402	151	31
169	552	0.968116	2.76388	178	35
170	1204	0.99177	1.912832	162	32
172	2010	0.993946	0.610072	155	36
173	1454	0.987296	1.893132	316	51
174	1056	0.979651	2.949199	335	53
175	824	0.95098	2.845177	368	46
176	851	0.96729	2.039242	202	47
177	778	0.985507	2.23077	221	25
178	2142	0.998093	0.359619	52	13

The coverage was estimated by a Good's estimator index. npshannon: non parametric index of Shannon. OTU classification was done with a treshold of 97% identity. doi:10.1371/journal.pone.0102649.t001

the abundance of each constituent bacterial genus). Accordingly, we analyzed the taxonomic structure of skin bacterial community using 16S tagged-amplicon pyrosequencing in 86 F2 fish progenies obtained in the previous work of Sauvage *et al.* (2012). Interindividual variation in abundance of each bacterial strain detected in host microbiota was afterwards projected on the results of the genetic linkage map in order to identify quantitative trait loci (QTL) of specific skin bacterial OTUs.

### **Materials and Methods**

# Ethics statement

All fish were reared and the experiment conducted strictly following guidelines required by the "Comité de Protection des Animaux de l'Université Laval (CPAUL, http://www.vrr.ulaval. ca/deontologie/cpa/index.html?accueil). The CPAUL reviewed and approved all experimental procedures used in this study.

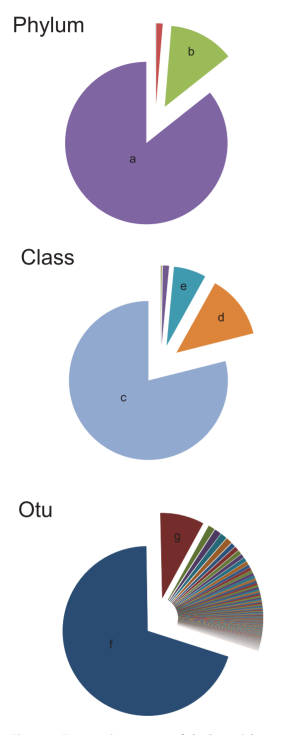


Figure 1. Taxonomic structure of the bacterial community of fish skin mucus at three different taxonomic levels: Phylum, Classl and OTU. a) Proteobacteria; b) Bacteroidetes, c) Alphaproteobacteria, d) Flavobacteria, e) Gammaproteobacteria, f) OTU 50 (*Methylobacterium rhodesianum*), g) OTU 36 (*Flavobacterium psychrophilum*).

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## Fish sampling

The study population was generated using two different populations of brook charr. The first one (D) is a domestic population used in aquaculture for more than 100 years

(Pisciculture de la Jacques Cartier - Cap-Santé, Québec, Canada). The other population (L), is an anadromous population from the Laval River near Forestville (North of the St. Lawrence River, Québec, Canada). Breeders from the L population were kept in captivity at the Station aquicole de l'ISMER (Québec, Canada,  $48^\circ31'N,\ 68^\circ28'W)$  under natural photoperiod and temperature. Crosses between 10 sires of each population (L and D; F<sub>0</sub> generation) with 10 dams (L and D) were performed to generate 10 full-sib outbred hybrid families (L×D -  $F_1$  generation). Subsequently, six F1 individuals were bi-parentally crossed to obtain three F2 families. The F2 family selected for the present study demonstrated the lowest post-hatch mortality rate (< 2%) and the greatest abundance. Fish were raised in the same tank and had fasted for 12 hours before sampling. Fish were measured  $(\text{mean} = 28.8 \pm 1.77 \text{ cm})$  and weighed  $(\text{mean} = 276.7 \pm 59.48 \text{ g})$  to calculate Fulton index [23].

Mucus was sampled using a sterile swab on the same area for all the fish [24]. We choose to sample the skin between the adipose fin and the caudal fin because this area was undisturbed by fish handling. Samples were stored in a sterile micro-centrifuge tube containing lysis buffer (Tris 50 mM, EDTA 40 mM, sucrose 0.75 g) and immediately stored in  $-80^{\circ}$ C until DNA extraction.

#### **DNA** extraction

DNA was extracted using a modified protocol of salt-extraction from Aljanabi & Martinez (1997). During the first lysis step, 22.6 µL of lysozyme (40 mg/mL) was added to the sample and incubated 45 minutes at 37°C. After this step, 22.6 µL of proteinase K (20 mg/µL) and 90 µL of SDS (10%) were added to the lysate and incubated at 55°C over night with agitation. The aqueous phase was transferred into a clean Eppendorf tube containing 600 µL of NaCl 6M, mixed and centrifuged 20 min at 14,800 rpm. The supernatant was transferred again into a clean Eppendorf tube containing 1 volume of cold isopropanol, mixed and stored 30 minutes at  $-20^{\circ}$ C. The mixture was centrifuged 20 minutes at 14,800 rpm and the supernatant was thrown away. The pellet was washed with cold ethanol 70%, air-dried and finally resuspended in 25 µL of sterile MilliQ H<sub>2</sub>O. Subsequently, DNA integrity and quantity were controlled using a Nanodrop instrument (ND-1000, Nanodrop).

#### Microbial 16S pyrosequencing

Each DNA sample, the 16S gene was PCR amplified using Takara Ex taq premix (Fisher). All PCR reactions were performed in a reaction volume of 50 µL containing 25 µL of premix Taq, 1 µM of each primer and sterile MilliQ H<sub>2</sub>O to up to 50 µL. A general reverse primer (R519) combined with B primer (Roche) was used for amplifications in combination with one of 86 uniquely tagged forward primers (F63-targeted) combined with A primer (Roche) (for primer sequences see [25,26]. The mean length of the amplified fragment was 450 bp. This procedure facilitates the parallel sequencing of thousands of samples on the same run and to reassign each reads to their respective samples. PCR conditions were applied as follows: after a denaturing step of 30 s at 98°C, samples were processed through 30 cycles consisting of 10 sec at 98°C, 30 sec at 55°C, and 30 sec at 72°C. The final extension step was done at 72°C for 4 min 30 sec. Following the amplification step, samples were purified using AMPure Beads (Beckman Coulter Genomics). Samples were adjusted to 100 µL with EB (Qiagen), 63 µL of beads were added. Samples were mixed and incubated for 5 min at RT. Using a Magnetic Particle Concentrator (MPC), the beads were pelleted against the wall of the tube and supernatant was removed. The beads were washed twice with 500  $\mu$ L of 70% ethanol and incubated for 30 sec each Table 2. Taxonomic variation observed between male and female.

Condition	N° OTU	Best Hit on Blast	Identity	Relative abundance
More abundant in female	82	Amaricoccus kaplicensis	98%	0.00037647
	115	Loktanella salsilacus	99%	0.00015338
	6	Methylovirgula ligni	91%	0.00026492
	163	Micrococcus antarcticus	99%	0.00061351
	88	Paracoccus yeei	94%	0.00019521
	75	Pseudochrobactrum kiredjianiae	99%	0.00050196
	128	Psychrobacter faecalis	99%	0.00100392
	131	Rheinheimera pacifica	99%	0.00057168
	7, 11	Singularimonas variicoloris	89%	0.00037647
	104	Sphingopyxis witflariensis	99%	0.00015338
	113	Stella humosa	100%	0.00018126
	142	Stenotrophomonas nitritireducens	99%	0.00022309
More abundant in male	186	Flavobacterium aquatile	97%	0.00029281
	190	Hyphomicrobium sulfonivorans	95%	0.00039041
	4	Methylosinus sporium	92%	0.00027887
	176	Polynucleobacter necessarius subsp. Asymbioticus	99%	0.00015338
	129	Pseudomonas hibiscicola	99%	0.00075294
	12	Pseudorhodoferax soli	85%	0.00018126
	140	Psychrobacter alimentarius	99%	0.00015338
	116	Rhodobacter blasticus	95%	0.00016732
	175	Simplicispira limi	99%	0.00034858

This table summarize the OTUs differentially abundant between skin mucus communities between male and female (Metastat using a FDR correction, p-value<0.01). doi:10.1371/journal.pone.0102649.t002

time. Supernatant was removed and beads were air dried for 5 min. Tubes were removed from the MPC and  $24 \ \mu$ L of EB were added. Samples were vortexed in order to suspend the beads. Finally, using the MPC, the beads were pelleted against the wall once more and supernatant was transferred to a new clean tube. Samples were quantified with Nanodrop before the amplification step. Amplicons were then quantified with Quant-iT PicoGreen dsDNA Assay Kit and mixed equally before being sent to the Plateforme d'Analyses Biomoléculaires (Institut de Biologie Intégrative et des Systèmes, Université Laval) for pyrosequencing on a 454 GS-FLX DNA Sequencer with the Titanium Chemistry (Roche), according to manufacturer's procedure.

#### 16S sequence analysis

All sequences are available on MG-RAST server (MG-RAST IDs: 4539915.3). The data were analyzed in two steps. First, CLC Genomics Workbench 3.1 (CLC Bio, Aarhus, Denmark CLC workbench BIO) was used to trim sequences for quality, recover and remove the primers' sequences and tags (minimum average quality score: 35 for a window of 50, number of differences to the primer sequence = 0, maximum number of differences to the barcode sequence = 0, number of ambiguous base calls = 0, maximum homopolymer length = 8). In a second step, preprocessing and analysis were performed using the MOTHUR software [27]. All datasets were checked for chimeras with the chimera slayer algorithm implemented in MOTHUR. Standardization of the different samples was done by using the zscore which calculates the normalized abundance as follows: normalized abundance = (abundance - mean) / standard deviation

[3,28,29]. An alternative method for normalization was also tested, which consisted of subsampling equal number of reads from each sample [30]. This method greatly reduced the numbers of sequences (224 sequences per samples), but gave the same results as zscore (Figure S2, Figure S3, Table S1). Therefore we preferred to keep zscore normalization to base our conclusions on a larger dataset. We used the Operational Taxonomic Unit-based method described by Costello et al. (2009) because it is not biased towards a predefined taxonomy. One index was retained to assess the quality of pyrosequencing: the sequence coverage index (Good's estimator). The sequence coverage index is a metric used to estimate the quality of the depth sequencing. All sequences were clustered into Operational Taxonomic Unit (OTU) using a 97% identity threshold and OTU were classified from phylum to genus using the program MOTHUR with the default settings. For all interesting OTUs (explaining the inter-individual variation or linked to a genetic region), we extracted all the sequences classified in that OTU and used the single best BLAST hit to identify the different species. Statistical differences in the abundance of each OTU were calculated with the software for metagenomic analysis (Metastats).

To visualize potential differences across host progenies in terms of mucus bacterial community structure, distances between those communities were computed using the Yue & Clayton measure of dissimilarity (Thetayc). This index developed by Yue and Clayton (2005) is a measure of dissimilarity between the structures of two communities, meaning that this calculator takes in account the abundances of each OTU. Then, Distances were represented using a dendrograms based on this index and statistical robustness of the dendrogram was determined by a Unifrac weighted test.

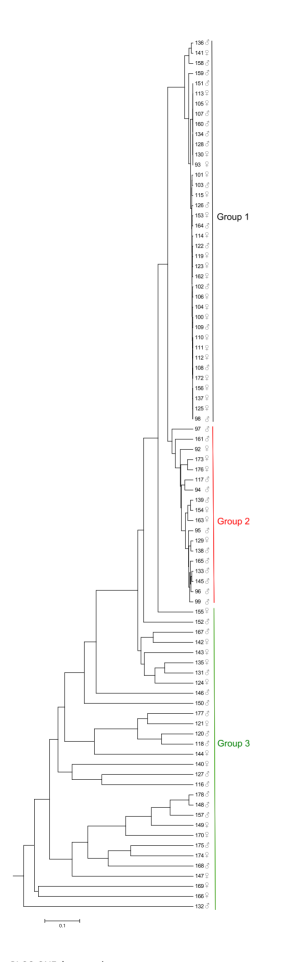


Figure 2. Dendrogram analysis based upon ThetaYC index of bacteria found on the skin of the 86 brook charr individuals. Groups are defined with the Weighted Unifrac distance. The first and second groups are composed of closely related bacterial communities. The third group is an assemblage of dissimilar communities and is considered as an outgroup. The most differentiated groups are groups 1 and 3 (Unifrac Score: 0.710811, p<0.0010) followed by the distance between groups 2 and 3 (Unifrac Score: 0.685361, p<0.0010), and the distance between groups 1 and 2 (Unifrac Score: 0.401674, p<0.0010). doi:10.1371/journal.pone.0102649.g002

This test allows determining whether any of the samples have a significantly different structure than the other samples. A random (Monte Carlo) permutation test was performed to test whether or not the distance between two communities was greater than expected by chance alone.

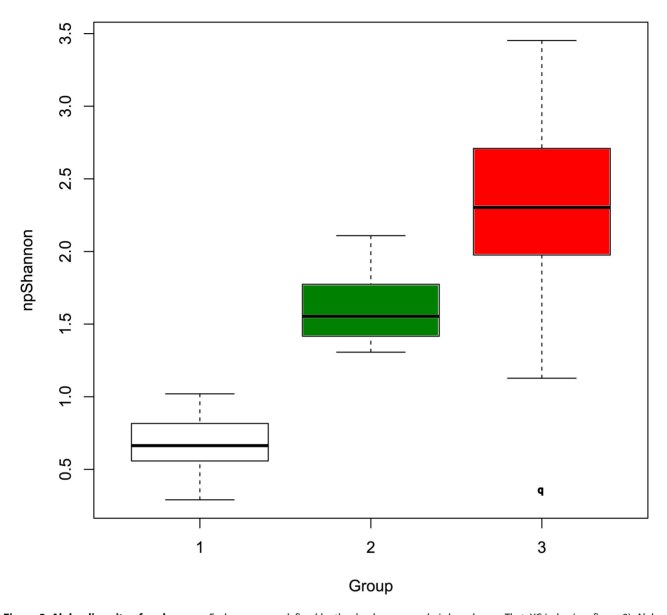
To visualize the distances between communities, we also applied a Principal Coordinate Analysis (PCoA) based on the Yue & Clayton measure of dissimilarity (Thetayc) using an eigenvectorbased approach to represent multidimensional data in as few dimensions as possible. This method allows determining which OTUs are responsible for shifting the samples along the PCoA axes by measuring the correlation of the relative abundance of each OTU clustered in genera with the axes of the PCoA dataset. Statistical differences in the abundance of each OTU was calculated to highlight which genera were the causative agents of the differentiation between groups obtained in the dendrogram based upon the TethaYC index [31]. All statistical analyses and graphics were carried out in the R environment (http://www.rproject.org).

## QTL detection

QTL analysis was carried out using the [R] package J/qtl [v. 1.3.1, August 2012, http://churchill.jax.org/software/jqtl.shtml/]. QTL were projected on the consensus brook charr linkage map (see Sauvage et al., 2012) according to the following steps. (1) A single QTL analysis was performed using the Haley-Knott (HK) regression method (10,000 permutations) to reveal which LGs were carrying QTL. The most probable position of the QTL was defined at the position giving the largest logarithm of odds (LOD) score indicating the QTL was fixed. (2) A two QTL model based upon the Haley-Knott regression was used to refine the QTL detection across the genome with a resolution of 5 cM and eventually to detect two QTL on a single LG linked to a particular trait. (3) The best model fitting our data was used to compute the percent variance explained (PVE expressed in %) by the QTL. The chromosome-wide and the genome-wide thresholds were calculated for each LG using 10,000 permutations. The 1.5 LOD confidence intervals were determined for all analyses following the Bayesian method implemented in the "bayesint" function in R/qtl. The bayesint function calculates an approximate interval (end points around the maximum LOD) for a given chromosome using the genome scan output.

### Results

Mucus samples were obtained from 86 F2 fish progenies (44 males and 42 females) issued from the same family. A total of 87,940 high-quality, classifiable sequences were obtained with an average number of  $1022\pm540$  per sample, which were subsequently classified in OTUs [1]. All of these sequences were successfully clustered into 9520 OTUs with 97% identity. A final trimming step was performed to focus on the most abundant OTU (which are represented by at least 10 sequences for the whole project) and result in a dataset containing 71,719 sequences (81% of the initial data set) clustered in 192 OTUs. The depth of



# **Bacterial diversity**

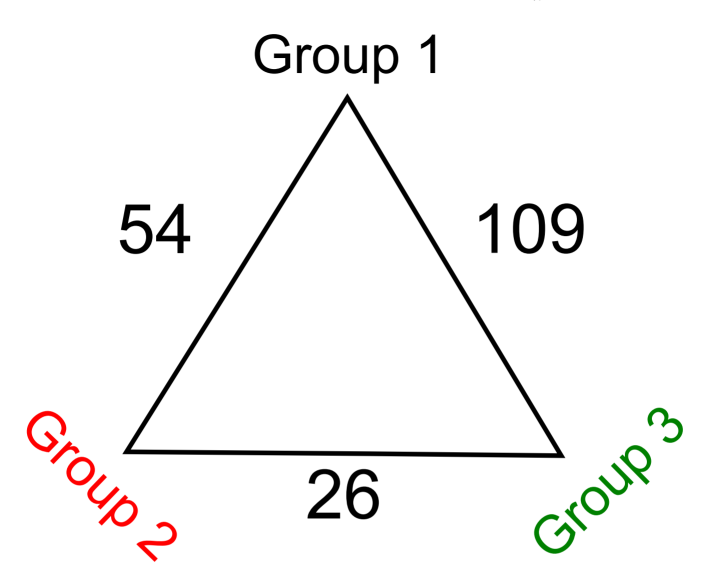
**Figure 3. Alpha diversity of each group.** Each group was defined by the dendrogram analysis based upon ThetaYC index (see figure 2). Alphadiversity was calculated by the non-parametric index of Shannon. doi:10.1371/journal.pone.0102649.g003

sequencing and the coverage were estimated using the Good's estimator index; the coverage found was always higher than 90% except for one sample (#127), which exhibited 76% of coverage (Table 1, Figure S1). The number of OTUs per sample was not equally distributed and the mean number was  $156.5\pm81.7$  (ranging from 33 to 368). The alpha-diversity was estimated by calculating the non-parametric index of Shannon (npShannon) because of the non-equally distributed abundance of each OTU in the samples. The npShannon index ranged from 0.29 to 3.45, with a mean index of  $1.43\pm0.83$ .

Phylum, class, and OTU composition of all samples were represented in the Figure 1. Most of the OTUs belong to the *Proteobacteria* phylum (88.7%) and the *Bacteroidetes* phyla (9.7%).

At the class level, most of the OTUs were classified as *Alpha-proteobacteria* (78.9%), *Gamma-proteobacteria* (6.5%) and *Flavo-bacteria* (13%). At the OTU level, the most abundant sequence was identified via BLAST as *M. rhodesianum* (69.8%) followed by *F. psychrophilum* (8%) (Figure 1 and Table S1).

Male and female brook charr harbored slightly different microbial communities. The Unifrac weighted distance between samples from males and females was statistically significant indicating that structures of bacterial communities were differentiated according to the sex of the host (Unifrac Score: 0.241695; p-value <0.001). However, there was not a sex difference in the two most abundant species, as this difference was based on 22 OTUs



**Figure 4. Descriptive analysis of the Metastats.** Numbers indicates the numbers of OTUs differentially present in the different groups of individuals. For details on the identities of the OTUs, please refer to the table 3; 4 and 5. doi:10.1371/journal.pone.0102649.q004

classified on 20 genera and 21 species, all of them being less abundant than 0.1% (Table 2).

In terms of community structure, clustering based on ThetaYC index clearly defined two groups of samples, whilst a third group was less well defined and composed of highly variable communities (Figure 2). There was little variation among samples in the first and second group (group 1 and group 2). The third group (group 3) was an assemblage of dissimilar communities and is considered as an external group. The variation of the alpha-diversity visualized by the npShannon index (Figure 3) indicated that alpha-diversity increases from group 1 to group 3. Furthermore, the bacterial communities within the third group were more diverse than groups 1 and 2. The same clustering was found with the normalization by subsampling (Figure S2, Figure S3, table S1). The difference between each group was greater than the difference between males and females (Figure 2). The most differentiated groups were groups 1 and 3 (Unifrac Score: 0.710811, p-value < 0.001) followed by the distance between groups 2 and 3 (Unifrac Score: 0.685361, p-value <0.001), and the distance between groups 1 and 2 (Unifrac Score: 0.401674, p-value <0.001). This was correlated with the numbers of OTU that were differentially abundant between those 3 groups (Figure 4). The divergence between groups 1 and 2 was based upon 54 OTU classified in 26 genera and 41 species (Table 3). The microbiota of the first group was dominated by *M. rhodesianum*, and host individuals from this group were tightly clustered. The second group was more diverse, dominated by *Flavobacterium* and had a lower abundance of *M. rhodesianum*. For groups 1 and 3, the difference was based upon 109 OTU classified in 45 genera and 70 species (Table 4). Finally, 26 OTU were differently abundant between group 2 and 3 and classified in 17 genera and 22 species (Table 5).

PCoA analysis displayed the same patterns of divergence between the three groups (Figure 5). Each group found in the Unifrac analysis was highlighted in the PCoA. To understand which OTUs were responsible for the differentiation of the cluster on the PCoA axes 1 and 2, individual correlation coefficients were calculated. Three OTUs were highly correlated with the two axes ( $\sigma$ >0.6); OTU 36, 50 and 149. Those OTU belonged to two species: *Flavobacterium psychrophilum* (OTU 36) and *M. Rhodesianum* (OTU 50 and 149). The negative correlation 
 Table 3. Detailed information related to the OTU differentially abundant between individuals belonging to group 1 and group 2.

Ν° ΟΤU	mean abundance in group 1	mean abundance in group 2	Best hit on Blast
36	0.00484	0.037926	Flavobacterium psychrophilum
122	0.000778	0.010066	Pseudomonas cedrina subsp. Fulgida
37	0.000471	0.006593	Flavobacterium terrigena
58	0.0014	0.006629	Methylocella palustris
128	0	0.004734	Psychrobacter faecalis
9	0.000531	0.004942	Pseudorhodobacter ferrugineus
90	0	0.004394	Hyphomicrobium sulfonivorans
39	0.000153	0.003833	Lysobacter capsici
70	0.000831	0.0043	Methylobacterium salsuginis
29	0.000028	0.003344	Methylobacterium adhaesivum
57	0.000518	0.003772	Rhodobacter blasticus
127	0.000218	0.003382	Acinetobacter haemolyticus
21	0.000117	0.002977	Flavobacterium aquatile
74	0	0.002743	Paracoccus yeei
52	0.000331	0.002911	Sphingopyxis alaskensis
126	0.000969	0.003524	Pseudomonas xanthomarina
89	0	0.002528	Sphingomonas dokdonensis
27	0.000113	0.001814	Sphingopyxis chilensis
63	0.00071	0.002374	Paracoccus haeundaensis
141	0.000156	0.001642	Acinetobacter johnsonii
84	0.000079	0.001484	Pseudorhodobacter ferrugineus
38	0.000063	0.001334	Sejongia jeonii
38	0.000014	0.001205	Acinetobacter junii
92	0	0.001099	Methylovirgula ligni
81	0.000248	0.00132	Ruegeria atlantica
;	0	0.001062	Methylosinus sporium
<b>i</b>	0	0.001033	Methylovirgula ligni
2	0	0.001029	Pseudorhodoferax soli
36	0.000282	0.00131	Methylobacterium rhodesianum
32	0.000125	0.001139	Methylobacterium zatmanii
22	0.000011	0.000871	Flavobacterium psychrophilum
107	0.000082	0.000932	Rhodobacter capsulatus
43	0.000258	0.001098	Flavobacterium aquatile
94	0.000073	0.000886	Rhodobacter sphaeroides
189	0.000043	0.000809	Methylomicrobium album
78	0.000047	0.000761	Rhodobacter capsulatus
1	0.000226	0.000929	Caulobacter leidyia
16	0.000220	0.000929	Flavobacterium psychrolimnae
101	0.00002	0.001111	Brevundimonas variabilis
154	0.00042	0.000728	Methylobacterium adhaesivum
154 32	0.000156		·
		0.000815	Amaricoccus kaplicensis
64 0	0.000176	0.000782	Methylopila capsulata
9	0.000154	0.000756	Brevundimonas variabilis
71	0	0.00054	Sphingomonas faeni
182	0.000046	0.000579	Flavobacterium aquatile
90	0.000282	0.000812	Methylobacterium adhaesivum
14	0.000094	0.000609	Flavobacterium psychrophilum
143	0	0.000513	Dokdonella koreensis

Table 3. Cont.					
N° OTU	mean abundance in group 1	mean abundance in group 2	Best hit on Blast		
97	0	0.00049	Hyphomicrobium facile subsp. Tolerans		
105	0	0.000359	Paracoccus haeundaensis		
137	0	0.000339	Acinetobacter johnsonii		
75	0.000558	0	Pseudochrobactrum kiredjianiae		
50	0.889343	0.677999	Methylobacterium rhodesianum		

Differentially abundancy were calculated by using Metastats (using a FDR correction, p<0.01). doi:10.1371/journal.pone.0102649.t003

between *Methylobacterium* and *Flavobacterium* was further confirmed by the co-occurrence analysis, which identified potential taxonomic interactions between genera. Two significant correlations between genera were detected: a positive correlation between *Maritimibacter* and *Micrococcus* ( $\sigma$ =0.69, p-value < 0.01) and a negative correlation between *Methylobacterium* and *Flavobacterium* ( $\sigma$ =-0.63, p-value <0.01). Furthermore, variance analysis of OTU 36' abundance showed that it was influenced by the Fulton index (Shapiro test for Normality: pvalue = 0.07, Linear model: p-value = 0.02741, F=5.0388, Rsq=-0.1496). A negative relationship between Fulton index and the OTU 36 was observed (Figure 6).

In the F2 fish progeny, three significant QTLs (at the genomewide level) were identified on two linkage groups (LG 11 and LG 16). One major QTL per strain was detected respectively for Lysobacter, Rheinheimera and Methylobacterium counts (LOD score = 9.89, 7.46 and 3.48 respectively). For each of these traits, the total PVE (percent variance explained) of the QTL were estimated to 17.01%, 31.05% and 41.1% for Methylobacterium, Rheinheimera and Lysobacter respectively. The most probable positions of these QTL, their respective 95% CIs, the closest linked molecular markers (one per QTL) as well as additive and dominance effects are presented in Table 6.

### Discussion

I.

Inter-individual variations in host microbiota has been well documented (e.g. [1]. Such variations occur even in hosts with identical genetic background, as observed in monozygotic twins [3] indicating that both genetic and environmental conditions play a role in the modulation of host microbiota. The gold standard forward genetics technique to identify areas of the genome that relate to certain phenotypes is to make a cross between genetically divergent individuals [32]. In this study, we focused on the structural variation of the skin microbiota (i.e abundance of each bacterial genus) of an F2 intercross originally generated from two genetically contrasted strains of brook charr. To our knowledge, this is the first study that identified host genomic regions associated with the abundance of specific microbiota strains in a non-model vertebrate.

The deep taxonomic analysis of the fish skin microbiota indicates that two phyla dominate: *Proteobacteria* (88.7%), followed by *Bacteroidetes* (9.7%). Those two phyla are mainly represented by a single OTU each; 50 and 36 respectively. OTU 50 is classified as M. *rhodesianum*. OTU 36 is classified as F.

*psychrophilum*, the causative agent of the cold water disease, a pervasive infectious disease in farmed fish [33,34]. This disease especially affects salmonids at early life-stages, and it is well documented that both stressful conditions, and injuries facilitate infection triggering [33,35]. According to the dendrogram (Figure 2), each individual's bacterial communities clustered into three groups. Similar clustering was also found with the second normalization method (subsampling of the same number of sequences for each samples). The same result was obtained with PCoA analysis and furthermore the Pearson correlation indicates that the genus Flavobacterium and Methylobacterium are negatively correlated. In the PCoA, two species (F. psychrophilum and M. rhodesianum) are correlated in opposite ways with the axis 1 which discriminate the three groups. We assume that two or more species which seem to be mutually exclusive are involved in a negative relationship [36]. All together those results indicate that the antagonistic relationship between those two dominant species is very likely playing a key role in shaping the structure of the individual's microbiota.

Because all fish progenies were reared in the same tank, and under the same environmental conditions since birth, environment had likely little influence on the phenotypic variation observed in this study. Yet, the individual's microbiota either clustered in one of two closely related groups, or formed divergent outliers. Sexspecific variations had little influence on individual clustering since both males and females were present in each defined group and there was no correlation between sex and groups. The difference between male and female is based on 22 OTUs which are less abundant than 0.1% of the community and are then classified as part of the rare biosphere. Previous studies observed similar individual variations of the microbiota in human or mouse [1,2,32].

Three OTUs classified into two species were responsible for the inter-individual differentiation; *M. rhodesianum and F. psychrophilum*. Interestingly, *Flavobacterium* and *Methylobacterium*, which are the most abundant genera, negatively co-occurred in the samples, indicating that the relationship between those two genera is based on competition. Furthermore, a strong negative correlation was found between the OTU 36 and 50, thus supporting a strong antagonistic relationship between the two species, *F. psychrophilum* and *M. rhodesianum*.

*M. rhodesianum* produces poly- $\beta$ -hydroxybutyrate, a polymer of short-chain fatty acid, known to inhibit the growth of pathogens like enterobacteria and *Vibrio sp.* [37,38,39,40,41,42]. Taken together, this suggests that a reduction of *M. rhodesianum* 

 Table 4. Detailed information related to the OTU differentially abundant between individuals belonging to group 1 and group 3.

υτο	mean abundance in group 1	mean abundance in group 3	Best hit on Blast
86	0.00484	0.15627	Flavobacterium psychrophilum
52	0.000331	0.052599	Sphingopyxis alaskensis
51	0.002582	0.045797	Sphingomonas paucimobilis
19	0.001237	0.034255	Pseudomonas peli
25	0.001567	0.030681	Acinetobacter lwoffii
20	0.002302	0.030821	Rheinheimera texasensis
3	0.00071	0.027857	Paracoccus haeundaensis
22	0.000778	0.018741	Pseudomonas cedrina subsp. Fulgida
1	0.003372	0.02093	Porphyrobacter dokdonensis
1	0.000117	0.011826	Flavobacterium aquatile
27	0.000218	0.00999	Acinetobacter haemolyticus
5	0	0.008105	Bradyrhizobium pachyrhizi
4	0.000176	0.007992	Methylopila capsulata
62	0.002112	0.009611	Knoellia sinensis
26	0.000969	0.008421	Pseudomonas xanthomarina
7	0.000518	0.007951	Rhodobacter blasticus
9	0.000531	0.007849	Pseudorhodobacter ferrugineus
1	0.000248	0.006693	Ruegeria atlantica
34	0.000245	0.00635	Acinetobacter Iwoffii
43	0	0.005914	Dokdonella koreensis
+3 29	0	0.005644	Pseudomonas hibiscicola
29	0	0.004908	Nevskia soli
)	0.000014	0.004576	Flavobacterium aquatile
3	0.000063	0.004547	Sejongia jeonii
3	0.0014	0.005784	Methylocella palustris
l	0	0.004232	Sphingomonas faeni
)	0.000259	0.004408	Rhodobacter capsulatus
8	0.000047	0.003657	Rhodobacter capsulatus
'	0.000471	0.004035	Flavobacterium terrigena
41	0.000156	0.003673	Acinetobacter johnsonii
56	0.000033	0.003487	Sphingopyxis alaskensis
9	0	0.00318	Flavobacterium psychrophilum
1	0.000095	0.00295	Pseudomonas peli
06	0.000106	0.002945	Bosea vestrisii
37	0	0.002615	Acinetobacter johnsonii
01	0.00042	0.002997	Brevundimonas variabilis
7	0	0.002498	Hyphomicrobium facile subsp. Toleran
39	0.000153	0.002639	Lysobacter capsici
57	0.00006	0.002488	Sphingomonas sanguinis
16	0	0.002339	Rhodobacter blasticus
77	0.000289	0.00255	Aquabacterium commune
i	0.000069	0.002236	Acinetobacter Iwoffii
32	0.000046	0.002172	Flavobacterium aquatile
7	0	0.002082	Flavobacterium psychrophilum
4	0.000079	0.00214	Pseudorhodobacter ferrugineus
64	0.000219	0.002164	Rhodococcus fascians
05	0	0.001788	Paracoccus haeundaensis
3	0.000258	0.002022	Flavobacterium aquatile
80	0.000017	0.001709	Acidovorax defluvii
2	0.000147	0.001809	Sphingopyxis witflariensis

# Table 4. Cont.

οτυ	mean abundance in group 1	mean abundance in group 3	Best hit on Blast
4	0	0.00161	Methylosinus sporium
145	0	0.001417	Acinetobacter johnsonii
5	0	0.001415	Magnetospirillum magnetotacticum
88	0.000036	0.00143	Amaricoccus veronensis
3	0	0.001358	Rhodobacter changlensis
38	0.000014	0.001365	Acinetobacter junii
0	0.000831	0.002052	Methylobacterium salsuginis
84	0	0.001156	Flavobacterium aquatile
22	0.000011	0.001144	Flavobacterium psychrophilum
12	0	0.001106	Sphingomonas sanguinis
18	0.000062	0.001143	Haematobacter missouriensis
3	0.000029	0.001093	Roseomonas mucosa
52	0.000028	0.001009	Rhodobacter blasticus
5	0.00005	0.000936	Sphingomonas sanguinis
48	0.000033	0.000915	Psychrobacter arcticus
72	0	0.000864	Sphingomonas sanguinis
46	0	0.000832	Pseudomonas mohnii
4	0.000073	0.000865	Rhodobacter sphaeroides
40	0	0.000775	Psychrobacter alimentarius
60	0.000061	0.000823	Sphingomonas faeni
3	0.000128	0.000877	Paracoccus aminovorans
2	0.000323	0.001071	Flavobacterium psychrophilum
86	0	0.000711	Flavobacterium aquatile
4	0.000094	0.000796	Flavobacterium psychrophilum
17	0.000029	0.000731	Sphingopyxis taejonensis
07	0.000082	0.000766	Rhodobacter capsulatus
2	0.000432	0.001098	Beijerinckia mobilis
- 04	0.000028	0.000693	Sphingopyxis witflariensis
13	0	0.000663	Stella humosa
15	0	0.000622	Loktanella salsilacus
08	0.000051	0.000647	Novosphingobium panipatense
6	0	0.000559	Acinetobacter johnsonii
8	0	0.00052	Flavobacterium psychrophilum
66	0	0.000508	Rhodococcus fascians
5	0.00064	0.001148	Methylobacterium organophilum
5 03	0.00084	0.000655	Sphingomonas ursincola
44	0.000215	0.000462	Acinetobacter Iwoffii
	0.00007	0.000462	Beijerinckia derxii subsp. Venezuelae
91			
9	0.000154	0.000498	Brevundimonas variabilis
78	0.000043	0.000378	Pseudorhodoferax soli
3	0.000017	0.000348	Flavobacterium psychrophilum
92	0	0.000328	Methylovirgula ligni
9	0.000058	0.000354	Flavobacterium aquatile
	0	0.00029	Methylosinus sporium
	0.000062	0.000245	Sphingomonas wittichii
69	0.000207	0.000287	Corynebacterium tuberculostearicun
11	0.000416	0.000202	Mycoplana bullata
,	0.000343	0	Singularimonas variicoloris
175	0.000348	0	Simplicispira limi

Table 5. Detailed information related to the OTU differentially abundant between individuals belonging to group 2 and group 3.

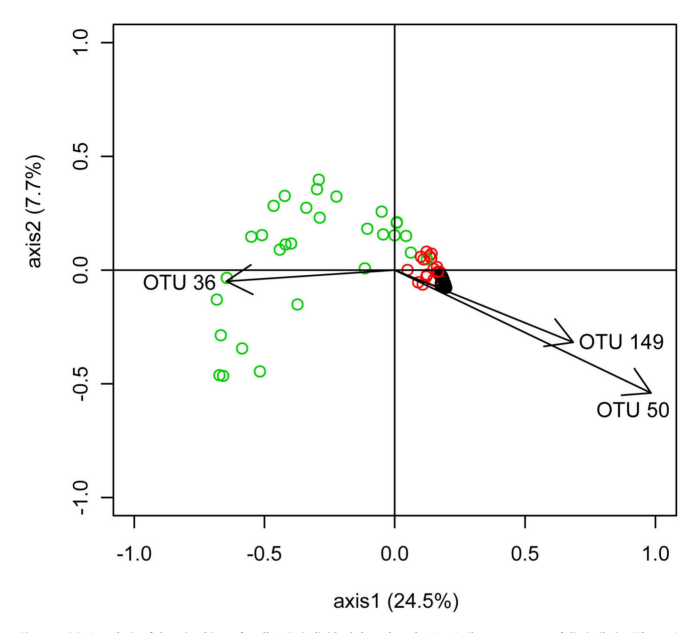
N° ΟΤU	mean abundance in group 2	mean abundance in group 3	Best hit on Blast
63	0.002374	0.027857	Paracoccus haeundaensis
55	0	0.008105	Bradyrhizobium pachyrhizi
54	0.000782	0.007992	Methylopila capsulata
134	0.000155	0.00635	Acinetobacter Iwoffii
129	0	0.005644	Pseudomonas hibiscicola
143	0.000513	0.005914	Dokdonella koreensis
124	0.000109	0.004908	Nevskia soli
40	0.000306	0.004576	Flavobacterium aquatile
123	0.000266	0.004124	Acinetobacter haemolyticus
71	0.00054	0.004232	Sphingomonas faeni
38	0.001334	0.004547	Sejongia jeonii
156	0.000306	0.003487	Sphingopyxis alaskensis
39	0.000144	0.00318	Flavobacterium psychrophilum
78	0.000761	0.003657	Rhodobacter capsulatus
137	0.000339	0.002615	Acinetobacter johnsonii
47	0.000087	0.002082	Flavobacterium psychrophilum
164	0.000236	0.002164	Rhodococcus fascians
4	0	0.00161	Methylosinus sporium
182	0.000579	0.002172	Flavobacterium aquatile
3	0	0.001358	Rhodobacter changlensis
184	0	0.001156	Flavobacterium aquatile
12	0.000252	0.001071	Flavobacterium psychrophilum
12	0.001029	0	Pseudorhodoferax soli
5	0.001033	0	Methylovirgula ligni
27	0.001814	0.000082	Sphingopyxis chilensis
50	0.677999	0.227483	Methylobacterium rhodesianum

Differentially abundancy were calculated by using Metastats (using a FDR correction, p<0.01). doi:10.1371/journal.pone.0102649.t005

abundance allows colonization or over-growth by other bacteria, some of which are pathogenic.

Evidently, a change in skin microbiota taxonomic structure favors opportunist bacteria and especially opportunistic pathogens like F. psychrophilum, Acinetobacter haemolyticus, Acinetobacter johnsonii and Acinetobacter junii, which have a higher abundance in group 2 compared with group 1 Such a disturbance in the microbiome homeostasis is called dysbiosis, and its occurrence enlightens the importance of the function of M. rhodesianum in controlling the balance between both other commensal bacteria and opportunistic pathogens. Furthermore, Flavobacterium psychrophilum was negatively correlated to both M. rhodesianum abundance and Fulton index. The Fulton index is a condition factor used as a proxy for fish health status. Therefore, it suggests that skin microbiota from the weakest host individuals were unable to prevent colonization by the opportunistic pathogen F. psychrophilum. Furthermore, skin microbiota taxonomic structure varies significantly across the F2 progeny. This result added to the fact that *i* the genetics of the host is the only variable in the experiment, and *ii* the finding of three QTLs associated to the abundance of bacterial genera with high PVE, suggest that host genotype influences the abundance of commensal strains e.g. Methylobacterium, which will regulate the abundance of F. psychrophilum.

We found three QTL associated with the abundance of three genera: Lysobacter, Rheinheimera and Methylobacterium, all of which were observed to provide antimicrobial compounds [39,40,41,42,43,44,45,46,47]. These finding strongly suggests that host genotype influences abundance of specific commensal strains, and possibly targets their recruitment. The most compelling evidence concerns the QTL associated with Methylobacterium (PVE = 17.01%): as previously described above, Methylobacterium is influential on the structure and the homeostasis of the microbiota. Its abundance is inversely correlated with those of the pathogen F. psychrophilum. The targeted recruitment of M. *rhodesianum* mediated by the host genotype is therefore a strategy to prevent pathogen growth via harnessing antagonistic relationship towards resources use [48]. To our knowledge, this is only the second study that identified QTL associated with microbial variation among individuals. A study on murine gut microbiota showed similar results, in which McKnite et al. (2012) found 5 OTL located on four chromosomes influencing the variation of

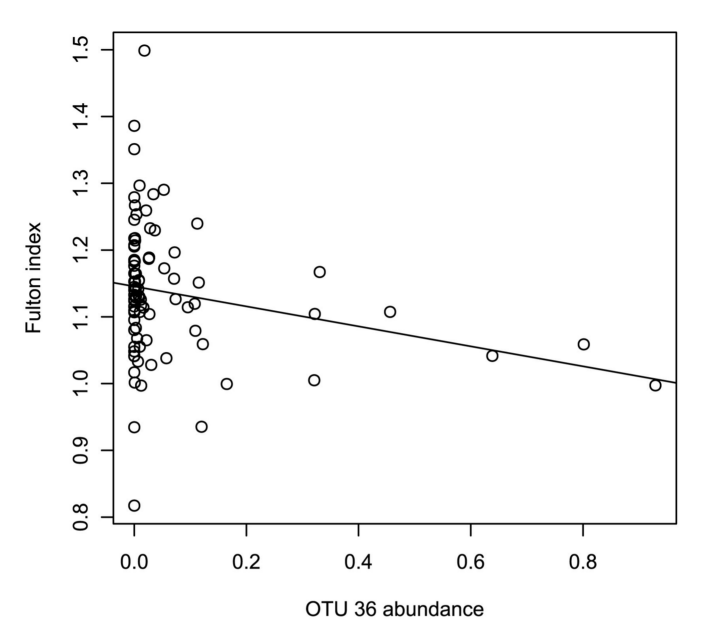


**Figure 5. PCoA analysis of the microbiome for all 86 F2 individuals based on the Yue & Clayton measure of dissimilarity (Thetayc).** Black circles represent individuals belonging to the group 1, red circles represent individuals belonging to the group 2 and green circles represent individuals belonging to the group 3. Arrows represent genus, which are significantly correlated with the axis. The first and second axes represented 24.5% and 7.7% of the variation respectively. The R-squared between the original distance matrix and the distance between the points in 2D PCoA space was 0.87.

doi:10.1371/journal.pone.0102649.g005

different taxa. The variance explained by their QTL is in the same range of our (20% to 27% of the variance explained for McKnite *et al.* (2012)). We also found QTLs with a major effect on the variance of genus abundances (PVE ranging for 17.02 to 41.1%). Linkage analysis on genus abundance data strongly evidenced the influence of host genetics on the modulation and/or recruitment for those genera. Therefore, brook charr immunity involves both specific commensal strains recruitment and nuclear gene expression, as previously evidenced in [49], those are under the control of the host genotype.

The study of the genetic architecture underlying the regulation of bacterial abundance further highlights the coevolutionary pattern between host, commensals, and pathogens. However, identifying genes located in the genomic regions (QTL) linked to the abundance of microbial partners will be far more challenging as the genome of brook charr is currently not fully sequenced and poorly annotated compared to the mouse genome. The markers surrounding the QTL regions may define regions to be deeply sequenced in future work to identify potential underlying genes and their associated functions. Evidently, those markers will be invaluable to conduct highly innovative genetic breeding programs targeting disease resistant host strains via the recruitment of highly resilient microbiota.



**Figure 6. Relationship between the Fulton Index and and the abundance of the OTU 36.** A linear regression is observed based on a linear model (Shapiro test for Normality: p-value = 0.07, Linear model: p-value = 0.02741, F = 5.0388, Rsq = 0.04536). doi:10.1371/journal.pone.0102649.g006

**Table 6.** Descriptive statistics, including the LOD score, the position, 95% CI, PVE (%), the associated P-value of each QTL linked to every phenotype related to bacteria counts trait (LOD, Log<sub>10</sub> of the odd ratio; 95% CI, 95% confidence interval; PVE, percent variance explained) [23].

Phenotype	Linkage Group	Pos (cM)	95% CI (cM)	LOD	<i>p</i> value (F)	PVE (in %)	Nearest marker
Lysobacter	11	62	56.2–68.2	9.89	0.000	41.1	sf003455
Rheinheimera	16	42.8	39.3-45.5	7.46	0.000	31.05	SFO-D91
Methylobacterium	16	39	34.5-43.5	3.48	0.001	17.02	SFO-D91

doi:10.1371/journal.pone.0102649.t006

#### **Supporting Information**

Figure S1 Rarefaction curves for each group defined by the dendrogram analysis based upon ThetaYC index (see figure 2). Each group curve reaches a plateau thus indicating the depth of sequencing is sufficient. (EPS)

Figure S2 Alpha diversity of each group normalized by subsampling of the same number of reads per sample. Each group was defined by the dendrogram analysis based upon ThetaYC index performed on the normalized dataset. Alphadiversity was calculated by the non-parametric index of Shannon. Statistical differences (represented by the asterisk) were calculated with a wilcoxon test with a correction for multiple testing (Holm, p-values  $<1.10^{-4}$ ). (EPS)

Figure S3 PCoA analysis of the microbiome for all 86  $F_2$  individuals. This PCoA was performed on a normalized dataset with the second method of normalization (subsampling of the same number of reads per sample). Black circles represent individuals belonging to the group 1, red circles represent individuals belonging to the group 2 and green circles represent individuals belonging to the group 3. Arrows represent genera, which are

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significantly correlated with the axis. The first and second axes represented 24.5% and 7.8% of the variation respectively. The Rsquared between the original distance matrix and the distance between the points in 2D PCoA space was 0.88. (EPS)

Table S1 Differentiation between the three groups of OTU calculated with an analysis of similarity (ANOSIM, statistic R). The q-values represent p-values corrected with Bonferroni and considered as significant when p<0.05. Note: The analysis was performed on a dataset normalized by two methods: zscore and subsampling. Both analyses gave the same results, each group being significantly different from the two others. (DOC)

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#### **Author Contributions**

Conceived and designed the experiments: ND CA LB SB CS. Performed the experiments: SB CS. Analyzed the data: SB CS. Contributed reagents/ materials/analysis tools: ND CA LB. Wrote the paper: ND CA LB SB CS.

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