### H.pylori Rhesus Microbiome Analysis

Noah Siegel

#### Data used in analysis

#### Library

```
suppressPackageStartupMessages({
  library(sjPlot)
  library(readxl)
  library(phyloseq)
  library(microbiome)
  library(DESeq2)
  library(qiime2R)
  library(tidyverse)
  library(tidyMicro)
  library(kableExtra)
  library(magrittr)
  library(ggpubr)
  library(microeco)
  library(tidytree)
  library(RColorBrewer)
  library(questionr)
  library(rmarkdown)
  library(stringr)
})
```

ASV counts the taxa table are linked below:

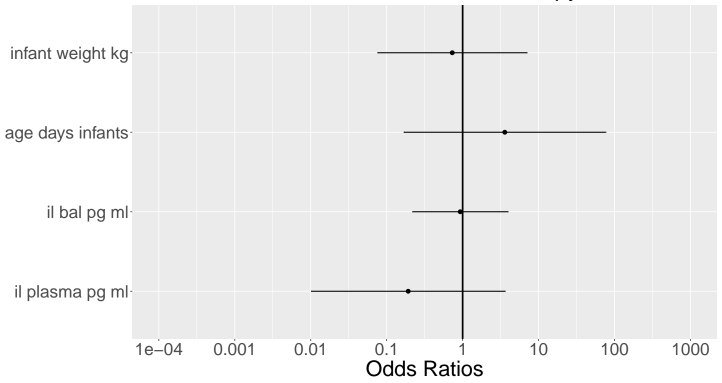
**ASV** counts

Taxa table

Table 1: H.pylori Metadata

SampleID	Treatment	site	Site_status	sex	log10_cfu.gm	IL8_Lavage	IL8_Plasma
LMiller_00458.BAL	H.pylori_(-)	BAL	H.pylori_(-)_BAL	Female	0.00	NA	463.376
LMiller_00458.Swab	H.pylori_(-)	Swab	H.pylori_(-)_Swab	Female	0.00	NA	463.376
LMiller_00459.BAL	H.pylori_(-)	BAL	H.pylori_(-)_BAL	Female	0.00	NA	258.416
LMiller_00459.Swab	H.pylori_(-)	Swab	H.pylori_(-)_Swab	Female	0.00	NA	258.416
$LMiller\_00461.BAL$	$H.pylori_(+)$	BAL	$H.pylori_(+)_BAL$	Female	7.66	NA	704.550
$LMiller\_00461.Swab$	$H.pylori_(+)$	Swab	$H.pylori_(+)_Swab$	Female	7.66	NA	704.550
$LMiller\_00462.BAL$	H.pylori_(-)	BAL	H.pylori_(-)_BAL	Female	0.00	19.104	1075.626
$LMiller\_00462.Swab$	H.pylori_(-)	Swab	H.pylori_(-)_Swab	Female	0.00	19.104	1075.626
$LMiller\_00463.BAL$	H.pylori_(-)	BAL	H.pylori_(-)_BAL	Female	0.00	11.590	423.590
$LMiller\_00466.BAL$	$H.pylori_(+)$	BAL	$H.pylori_(+)_BAL$	Female	3.81	11.201	642.148
$LMiller\_00466.Swab$	H.pylori_(+)	Swab	H.pylori_(+)_Swab	Female	3.81	11.201	642.148
$LMiller\_00467.BAL$	$H.pylori_(+)$	BAL	$H.pylori_(+)_BAL$	Female	6.09	15.511	425.782
$LMiller\_00467.Swab$	$H.pylori_(+)$	Swab	$H.pylori_(+)_Swab$	Female	6.09	15.511	425.782
$LMiller\_00468.BAL$	H.pylori_(-)	BAL	H.pylori_(-)_BAL	Female	0.00	15.511	660.911
$LMiller\_00468.Swab$	$H.pylori_{-}(-)$	Swab	H.pylori_(-)_Swab	Female	0.00	15.511	660.911
$LMiller\_00469.BAL$	$H.pylori_(+)$	BAL	$H.pylori_(+)_BAL$	Female	6.38	NA	1279.789
$LMiller\_00469.Swab$	$H.pylori_(+)$	Swab	$H.pylori_(+)_Swab$	Female	6.38	NA	1279.789
$LMiller\_00473.BAL$	H.pylori_(-)	BAL	H.pylori_(-)_BAL	Female	0.00	20.100	608.508
$LMiller\_00473.Swab$	H.pylori_(-)	Swab	H.pylori_(-)_Swab	Female	0.00	20.100	608.508
$LMiller\_00474.BAL$	$H.pylori_(-)$	BAL	H.pylori_(-)_BAL	Female	0.00	44.699	443.989
$LMiller\_00476.BAL$	H.pylori_(+)	BAL	H.pylori_(+)_BAL	Female	5.48	NA	476.431
LMiller 00476.Swab	$H.pylori_(+)$	Swab	H.pylori_(+)_Swab	Female	5.48	NA	476.431
$LMiller\_00477.BAL$	$H.pylori_(+)$	BAL	H.pylori_(+)_BAL	Female	5.81	NA	715.056
LMiller_00477.Swab	$H.pylori_(+)$	Swab	H.pylori_(+)_Swab	Female	5.81	NA	715.056
$LMiller\_00481.BAL$	H.pylori_(-)	BAL	H.pylori_(-)_BAL	Female	0.00	17.622	NA
$LMiller\_00481.Swab$	H.pylori_(-)	Swab	${\rm H.pylori}\_(\text{-})\_{\rm Swab}$	Female	0.00	17.622	NA

# Odds ratios of variables based on H. pylori status



	OR	2.5~%	97.5~%	p
(Intercept)	0.018	0.000	110.146	0.410
$infant\_weight\_kg$	0.243	0.000	7254.525	0.786
$age\_days\_infants$	1.042	0.955	1.177	0.413
$il\_bal\_pg.ml$	0.993	0.852	1.158	0.921
$il\_plasma\_pg.ml$	0.996	0.985	1.001	0.271

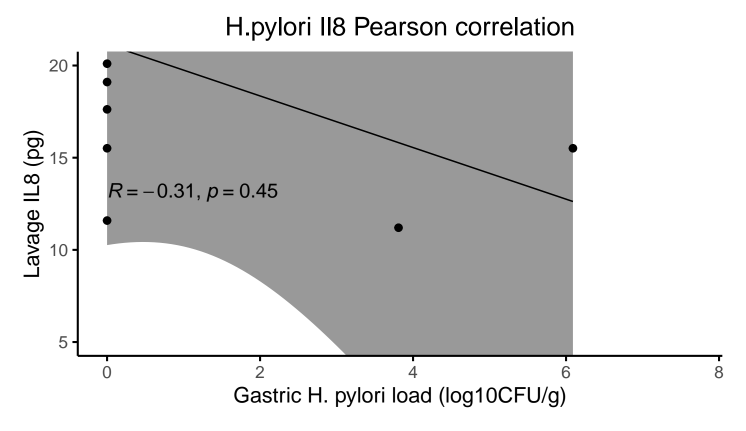


Figure 1: This plot only includes animals that had material sequenced and not all 25 animals from the study

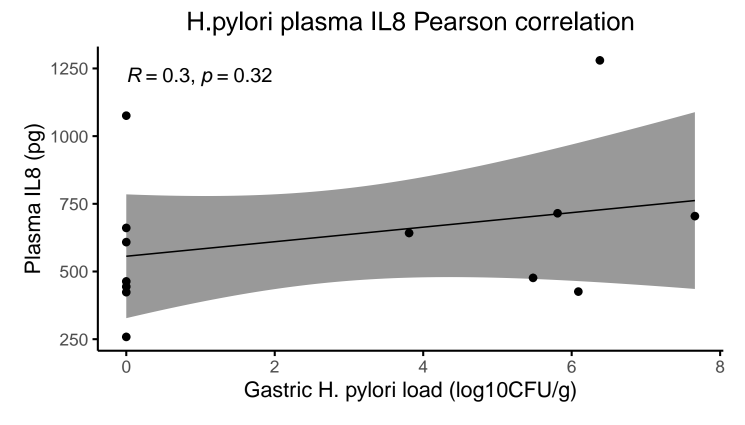
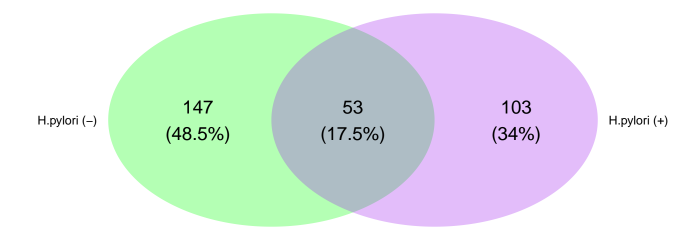
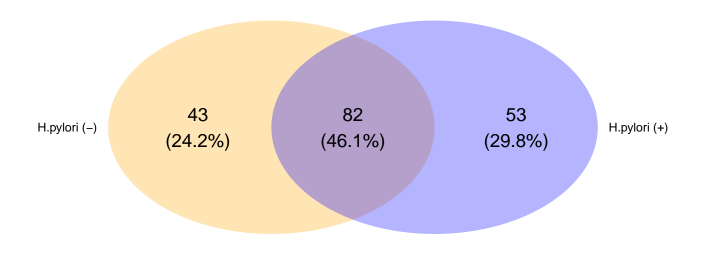


Figure 2: This plot only includes animals that had material sequenced and not all 25 animals from the study

# Overlapping taxa between H. pylori (+) and (-) in lavage

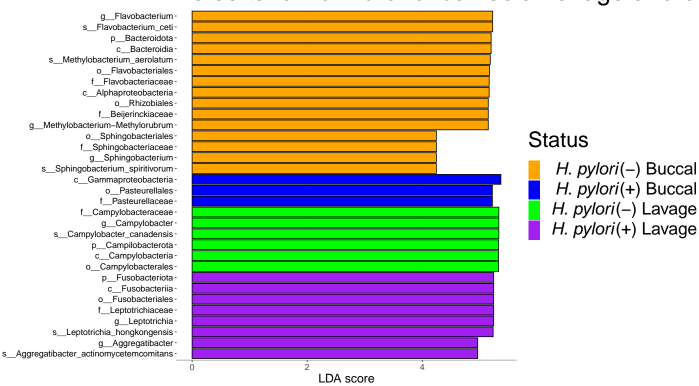


## Overlapping taxa between H. pylori (+) and (-) in buccal cavity

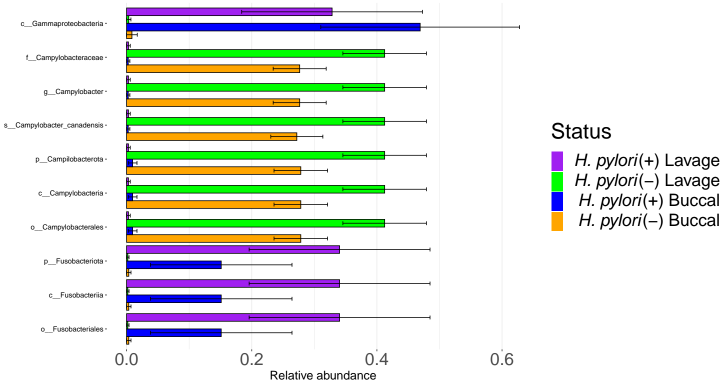


Lefse analysis and differential abundance for both sites

### Lefse taxa from bronchoalveolar lavage and or



## Lefse relative abundance from bronchoalveolar lavage



Lefse and differential abundance for bronchoalveolar lavage

-5.0

-25

#### Lefse Taxa from Bronchoalveolar Lavage p\_\_Proteobacteria c\_\_Alphaproteobacteria o\_\_Rhizobiales f\_\_Beijerinckiaceae g\_\_Methylobacterium-Methylorubrum s\_\_Methylobacterium\_aerolatum - Flavobacteriales f Flavobacteriaceae g\_\_Flavobacterium s\_\_Flavobacterium\_ceti-**Status** p\_\_Bacteroidota c\_\_Bacteroidia o\_\_Bacteroidales -H. pylori(-) Lavage o\_\_Sphingobacteriales -H. pylori(+) Lavage f\_Sphingobacteriaceae g\_\_Sphingobacterium s\_\_Sphingobacterium\_spiritivorumc\_\_Clostridia p\_\_Firmicutes p\_Campilobacterota c\_\_Campylobacteria o\_\_Campylobacterales f\_Campylobacteraceae g\_\_Campylobacters\_\_Campylobacter\_canadensis

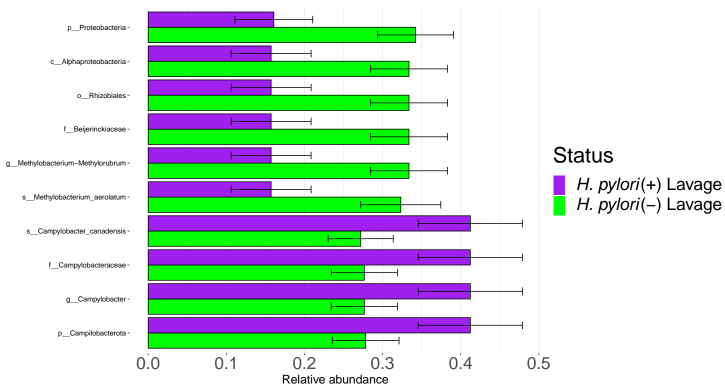
0.0

LDA score

2.5

5.0





Lefse and differential abundance for oral swabs

c\_\_Fusobacteriia o\_\_Fusobacteriales f\_\_Leptotrichiaceae g\_\_Leptotrichia -

-5.0

-2.5

#### p\_\_Proteobacteriac\_\_Gammaproteobacteriao\_\_Pasteurellales f\_\_Pasteurellaceae p\_\_Patescibacteriac\_\_Clostridia o\_\_Lachnospirales -**Status** f\_\_Lachnospiraceae p\_\_Firmicutes -H. pylori(-) Buccal H. pylori(+) Buccal g\_\_Aggregatibacters\_\_Aggregatibacter\_actinomycetemcomitans s\_\_Leptotrichia\_hongkongensisp\_\_Fusobacteriota-

Lefse taxa from oral swabs

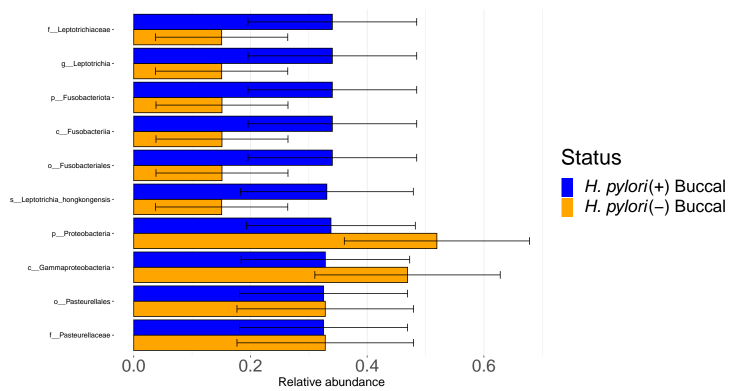
0.0

LDA score

2.5

5.0

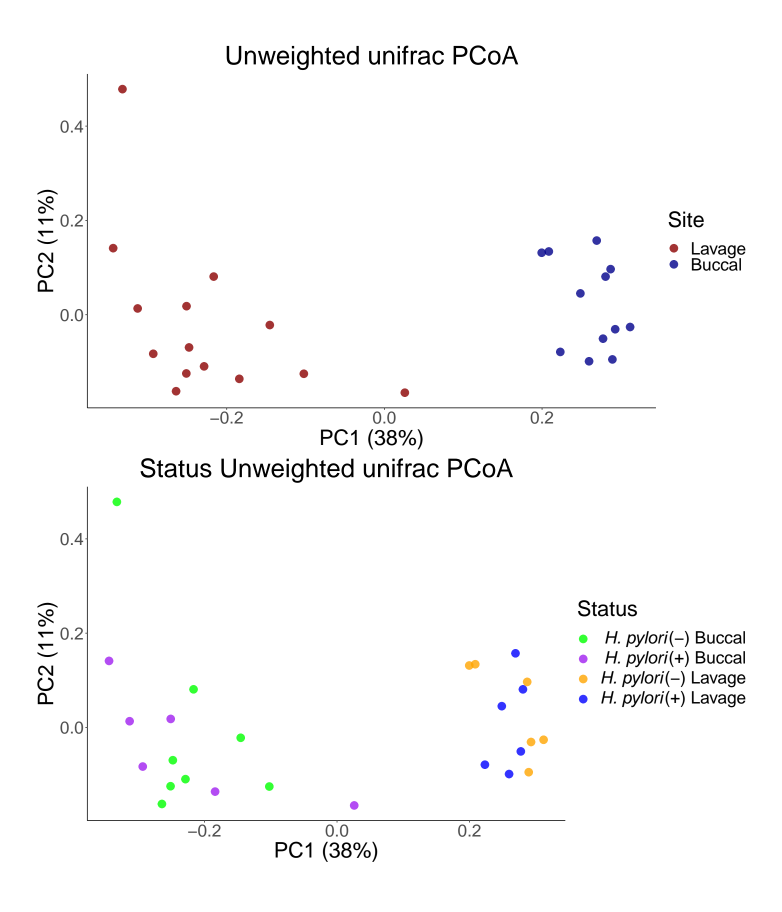
### Lefse relative abundance from oral swabs

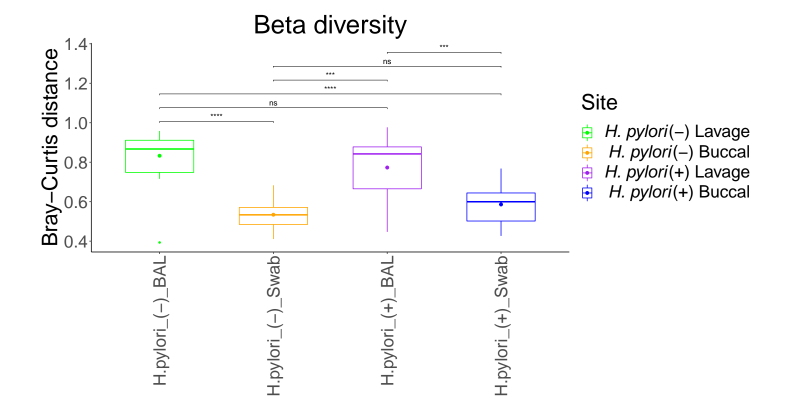


#### Beta diversity

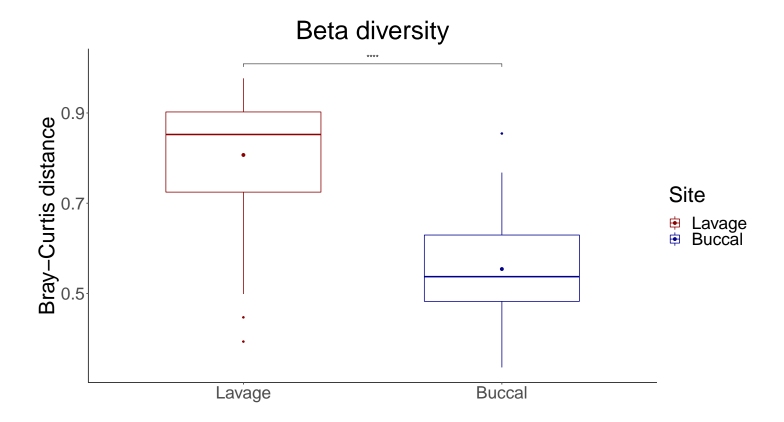
#### Unweighted unifrac PCoA plots

```
PC1
                   PC2
                              PC3
                                         PC4
                                                     PC5
                                                                PC6
                                                                           PC7
## 1 0.38311 0.1122317 0.08255574 0.05624649 0.04518453 0.04178459 0.03697168
            PC8
                      PC9
                               PC10
                                          PC11
                                                     PC12
                                                                PC13
## 1 0.03441675 0.0305229 0.0292508 0.02395872 0.0227312 0.01881898 0.01538496
          PC15
                     PC16
                                PC17
                                            PC18
                                                         PC19
                                                                     PC20
##
## 1 0.0140825 0.01226748 0.01155323 0.008776171 0.007488221 0.004824027
                        PC22
                                    PC23 PC24 PC25 PC26
##
            PC21
## 1 0.003528799 0.002630896 0.001679594
                                            0
```



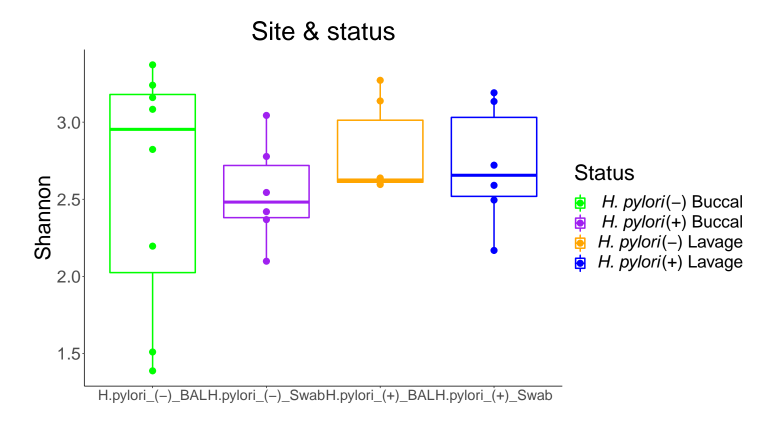


#### **Bray-Curtis**



#### Alpha Diversity

Shannon index by site and H. pylori status



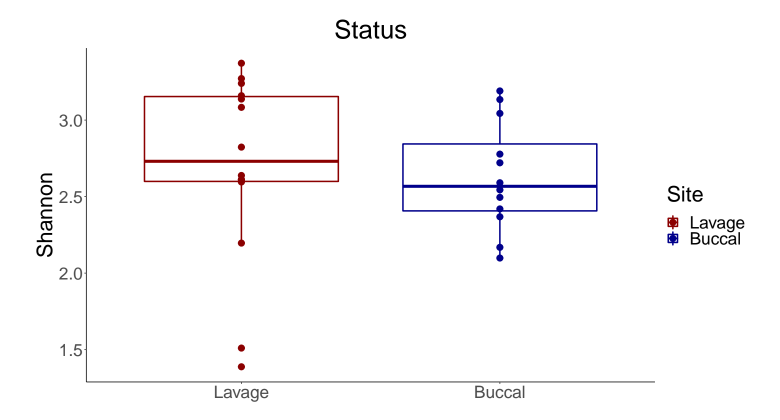
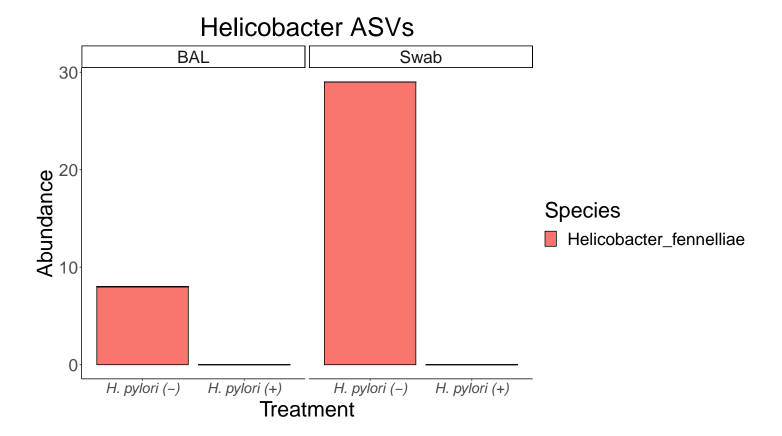


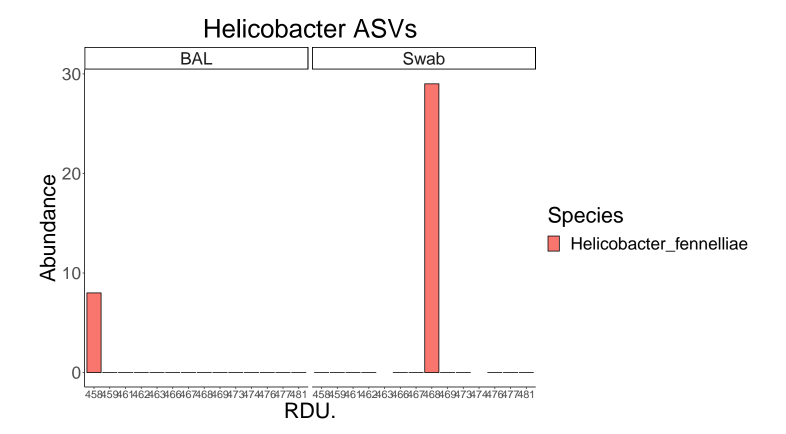
Table 2: All alpha diversity measurees

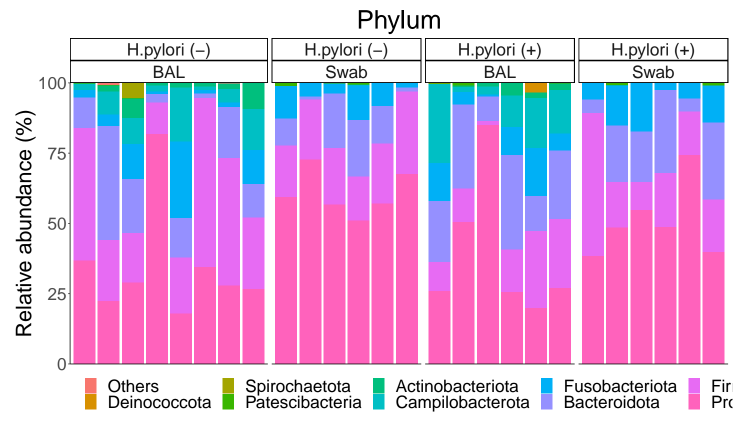
Groups	Measure	Test method	p.value	Significance
H.pylori_(-)_BAL vs H.pylori_(-)_Swab H.pylori_(-)_BAL vs H.pylori_(+)_BAL H.pylori_(-)_BAL vs H.pylori_(+)_Swab H.pylori_(-)_Swab vs H.pylori_(+)_BAL H.pylori_(-)_Swab vs H.pylori_(+)_Swab	Observed Observed Observed Observed	KW KW KW KW	0.220 0.651 0.081 0.262 0.520	
H.pylori_(+)_BAL vs H.pylori_(+)_Swab H.pylori_(-)_BAL vs H.pylori_(-)_Swab vs H.pylori_(+)_BAL vs H.pylori_(+)_Swab H.pylori_(-)_BAL vs H.pylori_(-)_Swab H.pylori_(-)_BAL vs H.pylori_(+)_BAL H.pylori_(-)_BAL vs H.pylori_(+)_Swab	Observed Observed Chao1 Chao1 Chao1	KW KW KW KW	0.109 0.202 0.220 0.651 0.081	
H.pylori_(-)_Swab vs H.pylori_(+)_BAL H.pylori_(-)_Swab vs H.pylori_(+)_Swab H.pylori_(+)_BAL vs H.pylori_(+)_Swab H.pylori_(-)_BAL vs H.pylori_(-)_Swab vs H.pylori_(+)_BAL vs H.pylori_(+)_Swab H.pylori_(-)_BAL vs H.pylori_(-)_Swab	Chao1 Chao1 Chao1 Chao1 ACE	KW KW KW KW	0.262 0.520 0.109 0.202 0.220	
H.pylori_(-)_BAL vs H.pylori_(+)_BAL H.pylori_(-)_BAL vs H.pylori_(+)_Swab H.pylori_(-)_Swab vs H.pylori_(+)_BAL H.pylori_(-)_Swab vs H.pylori_(+)_Swab H.pylori_(+)_BAL vs H.pylori_(+)_Swab	ACE ACE ACE ACE	KW KW KW KW	0.651 0.081 0.262 0.520 0.109	
H.pylori_(-)_BAL vs H.pylori_(-)_Swab vs H.pylori_(+)_BAL vs H.pylori_(+)_Swab H.pylori_(-)_BAL vs H.pylori_(-)_Swab H.pylori_(-)_BAL vs H.pylori_(+)_BAL H.pylori_(-)_BAL vs H.pylori_(+)_Swab H.pylori_(-)_Swab vs H.pylori_(+)_BAL	ACE Shannon Shannon Shannon Shannon	KW KW KW KW	0.202 0.439 1.000 0.796 0.109	
H.pylori_(-)_Swab vs H.pylori_(+)_Swab H.pylori_(+)_BAL vs H.pylori_(+)_Swab H.pylori_(-)_BAL vs H.pylori_(-)_Swab vs H.pylori_(+)_BAL vs H.pylori_(+)_Swab H.pylori_(-)_BAL vs H.pylori_(-)_Swab H.pylori_(-)_BAL vs H.pylori_(+)_BAL	Shannon Shannon Shannon Simpson Simpson	KW KW KW KW	0.337 0.423 0.540 0.519 0.699	
H.pylori_(-)_BAL vs H.pylori_(+)_Swab H.pylori_(-)_Swab vs H.pylori_(+)_BAL H.pylori_(-)_Swab vs H.pylori_(+)_Swab H.pylori_(+)_BAL vs H.pylori_(+)_Swab H.pylori_(-)_BAL vs H.pylori_(-)_Swab vs H.pylori_(+)_BAL vs H.pylori_(+)_Swab	Simpson Simpson Simpson Simpson	KW KW KW KW	0.699 0.109 0.423 0.262 0.493	
H.pylori_(-)_BAL vs H.pylori_(-)_Swab H.pylori_(-)_BAL vs H.pylori_(+)_BAL H.pylori_(-)_BAL vs H.pylori_(+)_Swab H.pylori_(-)_Swab vs H.pylori_(+)_BAL H.pylori_(-)_Swab vs H.pylori_(+)_Swab	InvSimpson InvSimpson InvSimpson InvSimpson InvSimpson	KW KW KW KW	0.519 0.699 0.699 0.109 0.423	
H.pylori_(+)_BAL vs H.pylori_(+)_Swab H.pylori_(-)_BAL vs H.pylori_(-)_Swab vs H.pylori_(+)_BAL vs H.pylori_(+)_Swab H.pylori_(-)_BAL vs H.pylori_(-)_Swab H.pylori_(-)_BAL vs H.pylori_(+)_BAL H.pylori_(-)_BAL vs H.pylori_(+)_Swab	InvSimpson InvSimpson Fisher Fisher Fisher	KW KW KW KW	0.262 0.493 0.606 0.699 0.796	
H.pylori_(-)_Swab vs H.pylori_(+)_BAL H.pylori_(-)_Swab vs H.pylori_(+)_Swab H.pylori_(+)_BAL vs H.pylori_(+)_Swab H.pylori_(-)_BAL vs H.pylori_(-)_Swab vs H.pylori_(+)_BAL vs H.pylori_(+)_Swab H.pylori_(-)_BAL vs H.pylori_(-)_Swab	Fisher Fisher Fisher Coverage	KW KW KW KW	0.631 0.631 1.000 0.927 NaN	NA
H.pylori_(-)_BAL vs H.pylori_(+)_BAL H.pylori_(-)_BAL vs H.pylori_(+)_Swab H.pylori_(-)_Swab vs H.pylori_(+)_BAL H.pylori_(-)_Swab vs H.pylori_(+)_Swab H.pylori_(+)_BAL vs H.pylori_(+)_Swab	Coverage Coverage Coverage Coverage	KW KW KW KW	NaN NaN NaN NaN NaN	NA NA NA NA
H.pylori_(-)_BAL vs H.pylori_(-)_Swab vs H.pylori_(+)_BAL vs H.pylori_(+)_Swab H.pylori_(-)_BAL vs H.pylori_(-)_Swab H.pylori_(-)_BAL vs H.pylori_(+)_BAL H.pylori_(-)_BAL vs H.pylori_(+)_Swab H.pylori_(-)_Swab vs H.pylori_(+)_BAL	Coverage PD PD PD PD	KW KW KW KW	NaN 0.366 0.699 0.439 0.631	NA
H.pylori_(-)_Swab vs H.pylori_(+)_Swab H.pylori_(+)_BAL vs H.pylori_(+)_Swab H.pylori_(-)_BAL vs H.pylori_(-)_Swab vs H.pylori_(+)_BAL vs H.pylori_(+)_Swab	PD PD PD	KW KW KW	0.749 $0.631$ $0.777$	

#### Helicobacter positive respiratory samples

Helicobacter positive samples separated by site







 ${\bf Figure~3:~Microbiota~Composition~at~Phylum~level}.$ 

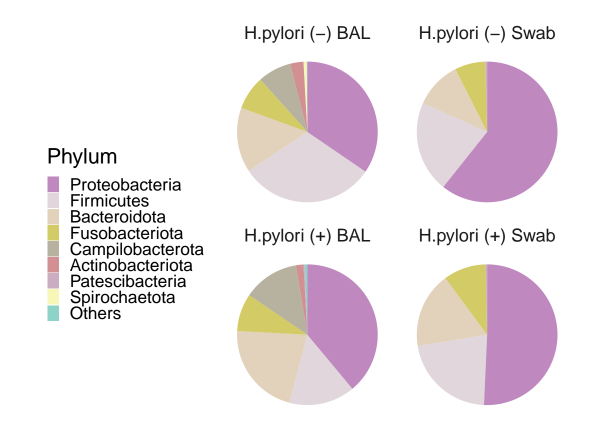


Figure 4: Microbiota Composition at Phylum level.

 Table 3: Phylum Average Relative Abundance

Taxonomy	Sample	Average Percent Abundance
Acidobacteriota Actinobacteriota Actinobacteriota Actinobacteriota Actinobacteriota	H.pylori (+) BAL H.pylori (-) BAL H.pylori (+) BAL H.pylori (+) Swab H.pylori (-) Swab	0.1% 3.1% 1.8% 0.1% 0%
Bacteroidota Bacteroidota Bacteroidota Bacteroidota Campilobacterota	H.pylori (+) BAL H.pylori (+) Swab H.pylori (-) BAL H.pylori (-) Swab H.pylori (+) BAL	21.7% $17.4%$ $14.9%$ $10.8%$ $12.9%$
Campilobacterota Campilobacterota Deinococcota Desulfobacterota Firmicutes	H.pylori (-) BAL H.pylori (-) Swab H.pylori (+) BAL H.pylori (-) BAL H.pylori (-) BAL	7.7% $0%$ $0.6%$ $0.1%$ $31.1%$
Firmicutes Firmicutes Firmicutes Fusobacteriota Fusobacteriota	H.pylori (+) Swab H.pylori (-) Swab H.pylori (+) BAL H.pylori (+) Swab H.pylori (+) BAL	21.7% $21%$ $15.2%$ $9.8%$ $8.6%$
Fusobacteriota Fusobacteriota Latescibacterota Patescibacteria Patescibacteria	H.pylori (-) BAL H.pylori (-) Swab H.pylori (-) BAL H.pylori (-) Swab H.pylori (+) Swab	7.8% $7%$ $0%$ $0.4%$ $0.3%$
Patescibacteria Patescibacteria Proteobacteria Proteobacteria Proteobacteria	H.pylori (+) BAL H.pylori (-) BAL H.pylori (-) Swab H.pylori (+) Swab H.pylori (+) BAL	0.3% $0%$ $60.7%$ $50.7%$ $38.9%$
Proteobacteria Spirochaetota	H.pylori (-) BAL H.pylori (-) BAL	$34.5\% \ 0.7\%$

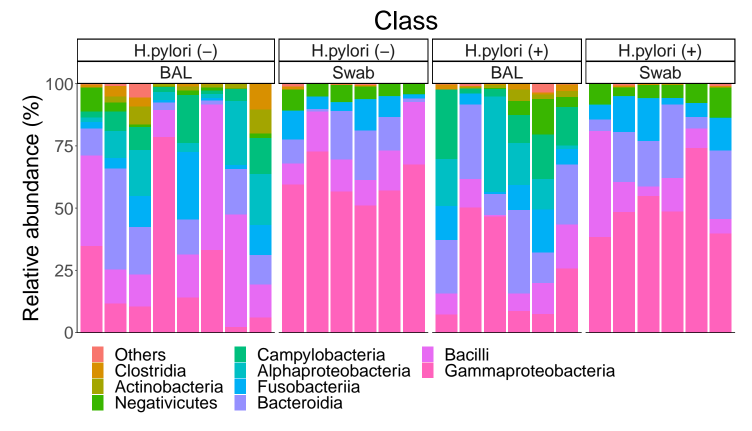


Figure 5: Microbiota Composition at Class level.

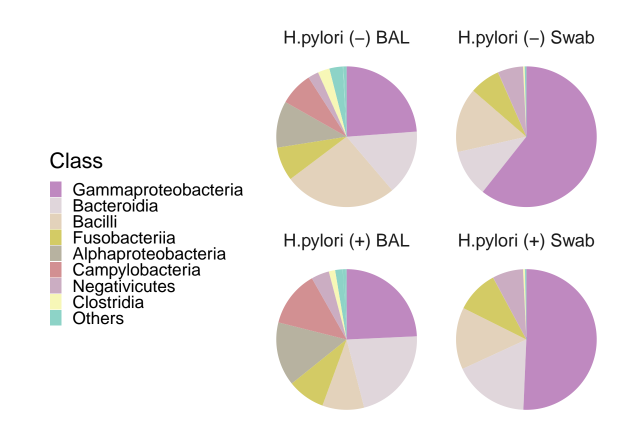


Figure 6: Microbiota Composition at Class level.

 Table 4: Class Average Relative Abundance

Taxonomy	Sample	Average Percent Abundance
Actinobacteria Actinobacteria Alphaproteobacteria Alphaproteobacteria Bacilli	H.pylori (-) BAL H.pylori (+) BAL H.pylori (+) BAL H.pylori (-) BAL H.pylori (-) BAL	3.1% 1.7% 14.7% 10.7% 26%
Bacilli Bacilli Bacilli Bacteroidia Bacteroidia	H.pylori (-) Swab H.pylori (+) Swab H.pylori (+) BAL H.pylori (+) BAL H.pylori (+) Swab	14.9% $14.2%$ $9.6%$ $21.7%$ $17.4%$
Bacteroidia Bacteroidia Campylobacteria Campylobacteria Clostridia	H.pylori (-) BAL H.pylori (-) Swab H.pylori (+) BAL H.pylori (-) BAL H.pylori (-) BAL	14.9% $10.8%$ $12.9%$ $7.7%$ $2.7%$
Clostridia Clostridia Clostridia Deinococci Fusobacteriia	H.pylori (+) BAL H.pylori (+) Swab H.pylori (-) Swab H.pylori (+) BAL H.pylori (+) Swab	1.5% $0.4%$ $0.4%$ $0.6%$ $9.8%$
Fusobacteriia Fusobacteriia Fusobacteriia Gammaproteobacteria Gammaproteobacteria	H.pylori (+) BAL H.pylori (-) BAL H.pylori (-) Swab H.pylori (-) Swab H.pylori (+) Swab	8.6% $7.8%$ $7%$ $60.7%$ $50.7%$
Gammaproteobacteria Gammaproteobacteria Negativicutes Negativicutes Negativicutes	H.pylori (+) BAL H.pylori (-) BAL H.pylori (+) Swab H.pylori (-) Swab H.pylori (+) BAL	24.3% $23.8%$ $7.1%$ $5.8%$ $4.1%$
Negativicutes Spirochaetia	H.pylori (-) BAL H.pylori (-) BAL	$2.4\% \\ 0.7\%$

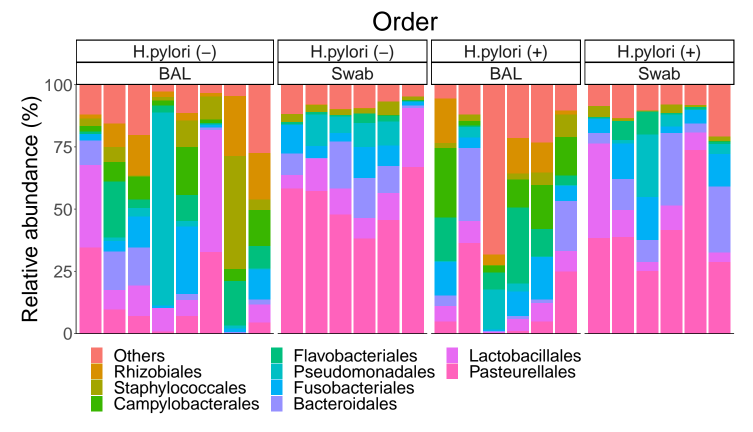


Figure 7: Microbiota Composition at Order level.

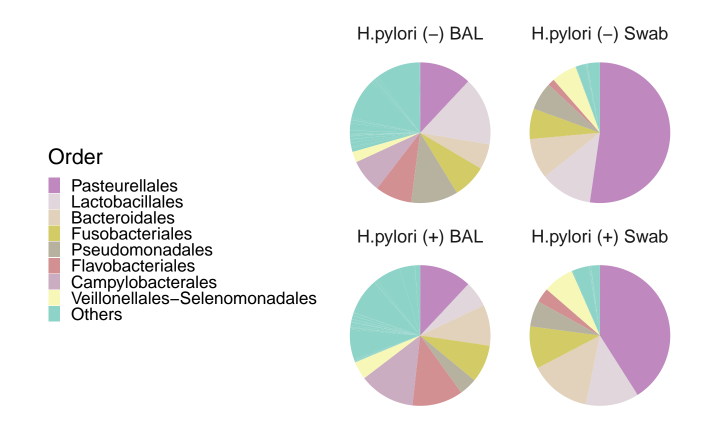


Figure 8: Microbiota Composition at Order level.

 Table 5: Order Average Relative Abundance

Taxonomy	Sample	Average Percent Abundance
Bacteroidales Bacteroidales Bacteroidales Bacteroidales Burkholderiales	H.pylori (+) Swab H.pylori (+) BAL H.pylori (-) Swab H.pylori (-) BAL H.pylori (+) BAL	14.1% 9.4% 9.3% 5.8% 6.8%
Campylobacterales Campylobacterales Flavobacteriales Flavobacteriales Fusobacteriales	H.pylori (+) BAL H.pylori (-) BAL H.pylori (+) BAL H.pylori (-) BAL H.pylori (+) Swab	12.9% 7.7% 11.7% 8.4% 9.8%
Fusobacteriales Fusobacteriales Fusobacteriales Lactobacillales Lactobacillales	H.pylori (+) BAL H.pylori (-) BAL H.pylori (-) Swab H.pylori (-) BAL H.pylori (+) Swab	8.6% $7.8%$ $15.7%$ $12.2%$
Lactobacillales Lactobacillales Pasteurellales Pasteurellales Pasteurellales	H.pylori (-) Swab H.pylori (+) BAL H.pylori (-) Swab H.pylori (+) Swab H.pylori (+) BAL	12% $6%$ $52.3%$ $41%$ $12%$
Pasteurellales Pseudomonadales Pseudomonadales Pseudomonadales Pseudomonadales	H.pylori (-) BAL H.pylori (-) BAL H.pylori (-) Swab H.pylori (+) Swab H.pylori (+) BAL	12% $10.8%$ $6.4%$ $5.9%$ $4.1%$
Rhizobiales Rhizobiales Sphingomonadales Staphylococcales Veillonellales-Selenomonadales	H.pylori (-) BAL H.pylori (+) BAL H.pylori (+) BAL H.pylori (-) BAL H.pylori (+) Swab	9.6% 8.3% 5.8% 10.1% 7.1%
Veillonellales-Selenomonadales Veillonellales-Selenomonadales	H.pylori (-) Swab H.pylori (+) BAL	5.8% 4.1%

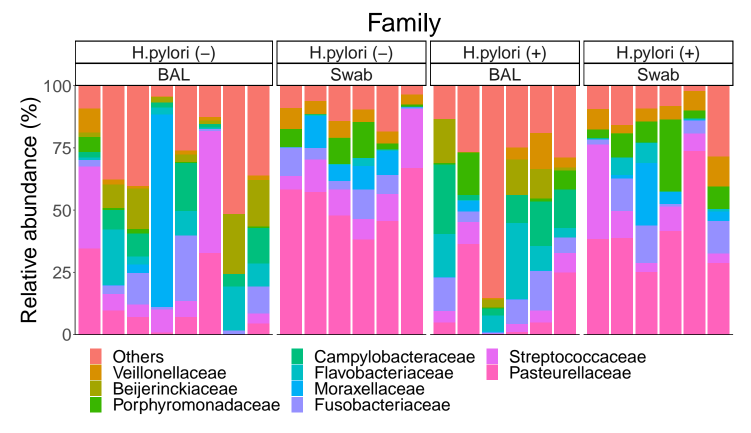


Figure 9: Microbiota Composition at Family level.

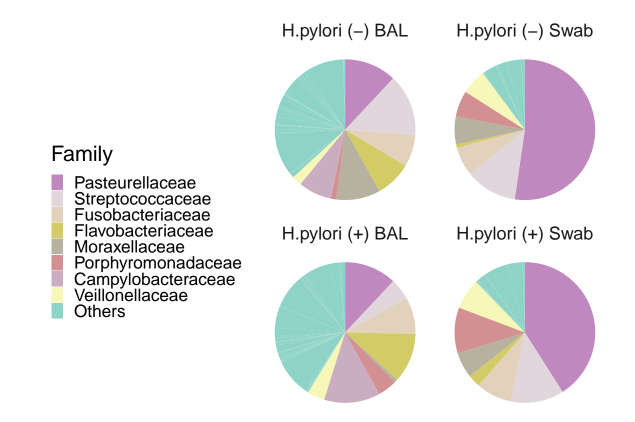


Figure 10: Microbiota Composition at Family level.

Table 6: Family Average Relative Abundance

Taxonomy	Sample	Average Percent Abundance
Beijerinckiaceae Beijerinckiaceae Campylobacteraceae Campylobacteraceae Flavobacteriaceae	H.pylori (-) BAL H.pylori (+) BAL H.pylori (+) BAL H.pylori (-) BAL H.pylori (+) BAL	9.5% 8.1% 12.9% 7.7% 11.5%
Flavobacteriaceae Fusobacteriaceae Fusobacteriaceae Fusobacteriaceae Fusobacteriaceae	H.pylori (-) BAL H.pylori (+) BAL H.pylori (+) Swab H.pylori (-) BAL H.pylori (-) Swab	8.3% $8.5%$ $8.2%$ $7.4%$ $6.6%$
Moraxellaceae Moraxellaceae Moraxellaceae Oxalobacteraceae Pasteurellaceae	H.pylori (-) BAL H.pylori (-) Swab H.pylori (+) Swab H.pylori (+) BAL H.pylori (-) Swab	10.1% $6.4%$ $5.9%$ $5%$ $52.3%$
Pasteurellaceae Pasteurellaceae Pasteurellaceae Porphyromonadaceae Porphyromonadaceae	H.pylori (+) Swab H.pylori (+) BAL H.pylori (-) BAL H.pylori (+) Swab H.pylori (-) Swab	41% $12%$ $12%$ $10.5%$ $5.9%$
Porphyromonadaceae Prevotellaceae Prevotellaceae Sphingomonadaceae Staphylococcaceae	H.pylori (+) BAL H.pylori (+) BAL H.pylori (-) BAL H.pylori (+) BAL H.pylori (-) BAL	4.4% $4.3%$ $4.1%$ $5.8%$ $8.4%$
Streptococcaceae Streptococcaceae Streptococcaceae Streptococcaceae Veillonellaceae	H.pylori (-) BAL H.pylori (+) Swab H.pylori (-) Swab H.pylori (+) BAL H.pylori (+) Swab	14.2% $12.2%$ $12%$ $4.9%$ $7%$
Veillonellaceae Veillonellaceae	H.pylori (-) Swab H.pylori (+) BAL	5.8% $4%$

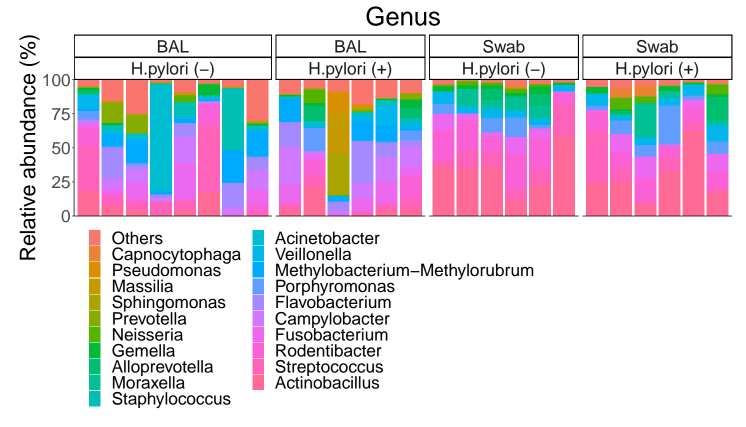


Figure 11: Microbiota Composition at Genus level.

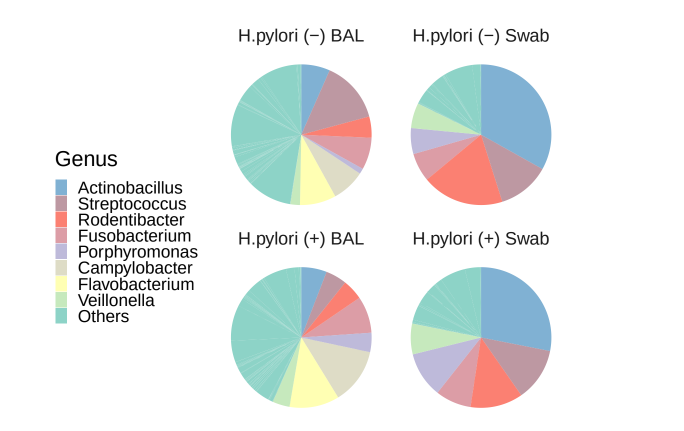


Figure 12: Microbiota Composition at Genus level.

 Table 7: Genus Average Relative Abundance

Taxonomy	Sample	Average Percent Abundance
Acinetobacter Actinobacillus Actinobacillus Actinobacillus Actinobacillus	H.pylori (-) BAL H.pylori (-) Swab H.pylori (+) Swab H.pylori (-) BAL H.pylori (+) BAL	10.1% 33.1% 28.1% 6.6% 5.9%
Campylobacter Campylobacter Flavobacterium Flavobacterium Fusobacterium	H.pylori (+) BAL H.pylori (-) BAL H.pylori (+) BAL H.pylori (-) BAL H.pylori (+) BAL	12.9% 7.7% 11.5% 8.3% 8.5%
Fusobacterium Fusobacterium Fusobacterium Massilia Methylobacterium-Methylorubrum	H.pylori (+) Swab H.pylori (-) BAL H.pylori (-) Swab H.pylori (+) BAL H.pylori (-) BAL	8.2% 7.4% 6.6% 4.9% 9.5%
Methylobacterium-Methylorubrum Moraxella Moraxella Porphyromonas Porphyromonas	H.pylori (+) BAL H.pylori (-) Swab H.pylori (+) Swab H.pylori (+) Swab H.pylori (-) Swab	8.1% 6.4% 5.9% 10.5% 5.9%
Rodentibacter Rodentibacter Rodentibacter Rodentibacter Sphingomonas	H.pylori (-) Swab H.pylori (+) Swab H.pylori (-) BAL H.pylori (+) BAL H.pylori (+) BAL	18.9% 12.1% 4.9% 4.7% 5.4%
Staphylococcus Streptococcus Streptococcus Streptococcus Streptococcus	H.pylori (-) BAL H.pylori (-) BAL H.pylori (+) Swab H.pylori (-) Swab H.pylori (+) BAL	8.4% 14.2% 12.2% 12% 4.9%
Veillonella Veillonella	H.pylori (+) Swab H.pylori (-) Swab	7% 5.8%

### Lavage Genus Abundance

Grouped by H. pylori status

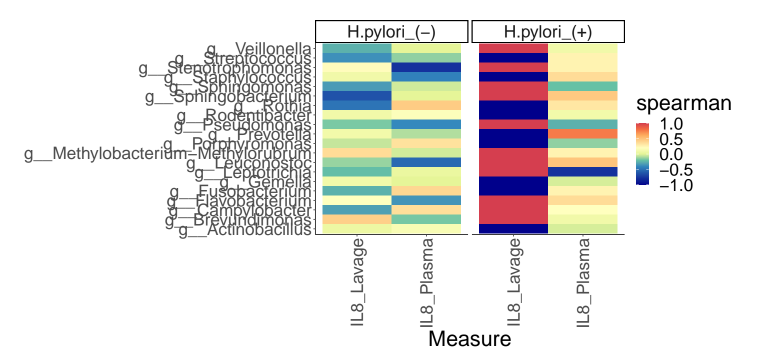


Figure 13: IL8 and lavage genus abundance Spearman correlations separated by H. pylori status

## Lavage Genus Abundance

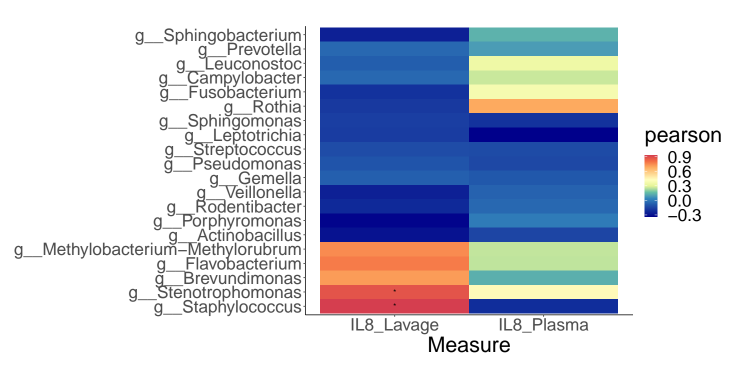


Figure 14: IL8 and lavage genus abundance Spearman correlations independent of H. pylori status

# Lavage Alpha Diversity

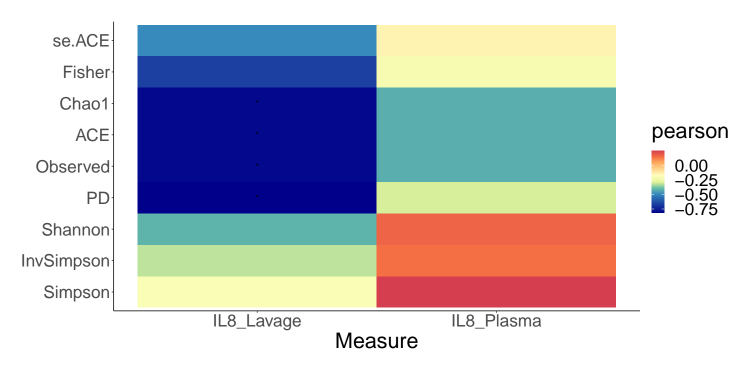


Figure 15: IL8 and lavage alpha diversity Spearman correlations inpendent of H. pylori status.

## **Buccal Cavity Genus Abundance**

Grouped by H. pylori status

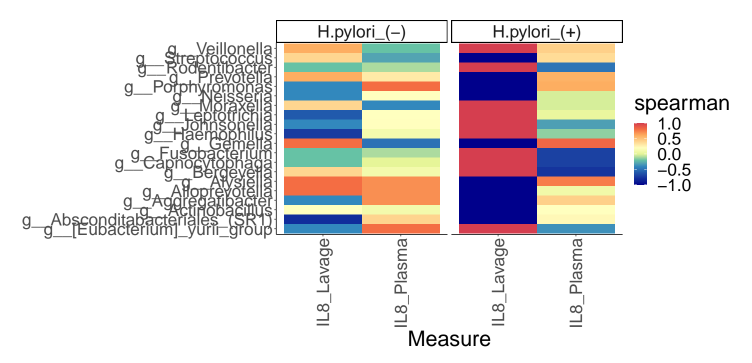


Figure 16: IL8 and Buccal genus abundance Spearman correlations separated by H. pylori status

## **Buccal Cavity Genus Abundance**

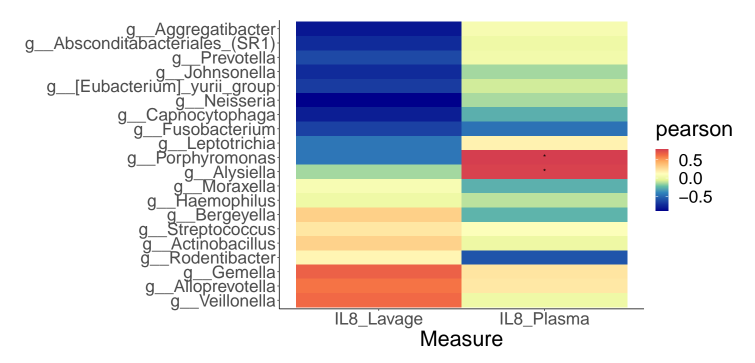


Figure 17: IL8 and Buccal genus abundance Spearman correlations independent of H. pylori status

# **Buccal Cavity Alpha Diversity**

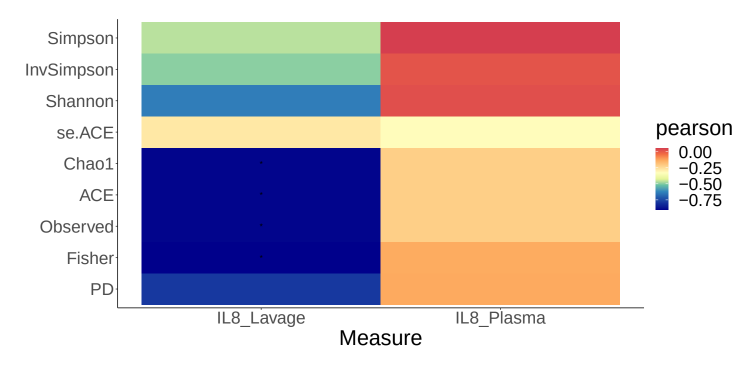


Figure 18: IL8 and Buccal alpha diversity Spearman correlations independent of H. pylori status

#### Record session information

```
## R version 4.1.3 (2022-03-10)
## Platform: x86_64-conda-linux-gnu (64-bit)
## Running under: Ubuntu 18.04.6 LTS
##
## Matrix products: default
## BLAS/LAPACK: /srv/conda/envs/notebook/lib/libopenblasp-r0.3.20.so
##
## locale:
##
   [1] LC_CTYPE=en_US.UTF-8
                                    LC_NUMERIC=C
##
   [3] LC_TIME=en_US.UTF-8
                                   LC_COLLATE=en_US.UTF-8
   [5] LC_MONETARY=en_US.UTF-8
                                    LC_MESSAGES=en_US.UTF-8
## [7] LC_PAPER=en_US.UTF-8
                                   LC_NAME=C
   [9] LC ADDRESS=C
                                   LC TELEPHONE=C
##
## [11] LC_MEASUREMENT=en_US.UTF-8 LC_IDENTIFICATION=C
##
## attached base packages:
## [1] stats4
                 stats
                           graphics grDevices utils
                                                          datasets methods
## [8] base
##
## other attached packages:
## [1] rmarkdown_2.14
                                     questionr_0.7.7
## [3] RColorBrewer_1.1-3
                                     tidytree_0.3.9
## [5] microeco_0.3.2
                                     ggpubr_0.4.0
##
   [7] magrittr_2.0.3
                                     kableExtra_1.3.4
## [9] tidyMicro_1.48
                                     forcats_0.5.1
## [11] stringr_1.4.0
                                     dplyr_1.0.9
## [13] purrr_0.3.4
                                     readr_2.1.2
## [15] tidyr_1.2.0
                                     tibble_3.1.7
                                     qiime2R_0.99.6
## [17] tidyverse_1.3.1
## [19] DESeq2 1.34.0
                                     SummarizedExperiment 1.24.0
## [21] Biobase_2.54.0
                                     MatrixGenerics_1.6.0
## [23] matrixStats 0.62.0
                                     GenomicRanges 1.46.1
## [25] GenomeInfoDb_1.30.0
                                     IRanges_2.28.0
## [27] S4Vectors 0.32.3
                                     BiocGenerics 0.40.0
## [29] microbiome_1.16.0
                                     ggplot2_3.3.6
## [31] phyloseq_1.38.0
                                     readxl 1.4.0
## [33] sjPlot_2.8.10
##
## loaded via a namespace (and not attached):
##
     [1] estimability_1.3
                                bit64_4.0.5
                                                        knitr_1.39
##
     [4] DelayedArray_0.20.0
                                 data.table_1.14.2
                                                        rpart_4.1.16
##
    [7] KEGGREST_1.34.0
                                RCurl_1.98-1.7
                                                        generics_0.1.2
##
   [10] timeSeries_3062.100
                                RSQLite_2.2.8
                                                        VGAM_1.1-7
##
   [13] bit_4.0.4
                                tzdb_0.3.0
                                                        webshot_0.5.3
##
   [16] xml2_1.3.3
                                 lubridate_1.8.0
                                                        httpuv_1.6.5
                                                        fBasics_3042.89.1
##
   [19] assertthat_0.2.1
                                xfun_0.31
##
    [22] hms 1.1.1
                                 evaluate 0.15
                                                        promises 1.2.0.1
##
   [25] fansi_1.0.3
                                dbplyr_2.2.0
                                                        igraph_1.3.1
##
   [28] DBI 1.1.2
                                geneplotter_1.72.0
                                                        htmlwidgets 1.5.4
   [31] ellipsis_0.3.2
                                backports_1.4.1
                                                        insight_0.17.1
##
##
   [34] permute 0.9-7
                                picante_1.8.2
                                                        annotate_1.72.0
##
   [37] deldir_1.0-6
                                vctrs_0.4.1
                                                        sjlabelled_1.2.0
##
   [40] abind_1.4-5
                                 cachem_1.0.6
                                                        withr_2.5.0
    [43] checkmate_2.1.0
                                 emmeans_1.7.4-1
##
                                                        vegan_2.6-2
##
    [46] svglite_2.1.0
                                 cluster_2.1.3
                                                        ape_5.6-2
##
   [49] lazyeval_0.2.2
                                 crayon_1.5.1
                                                        genefilter_1.76.0
##
   [52] pkgconfig_2.0.3
                                 {\tt zCompositions\_1.4.0-1 \ labeling\_0.4.2}
##
    [55] nlme_3.1-157
                                nnet_7.3-17
                                                        rlang_1.0.2
```

##		spatial_7.3-15	lifecycle_1.0.1	miniUI_0.1.1.1
##	[61]	modelr_0.1.8	randomForest_4.7-1.1	cellranger_1.1.0
##	[64]	datawizard_0.4.1	Matrix_1.4-1	carData_3.0-5
##	[67]	Rhdf5lib_1.16.0	boot_1.3-28	reprex_2.0.1
##	[70]	base64enc_0.1-3	png_0.1-7	viridisLite_0.4.0
##	[73]	stabledist_0.7-1	parameters_0.18.1	bitops_1.0-7
##	[76]	rhdf5filters_1.6.0	Biostrings_2.62.0	blob_1.2.3
##	[79]	GUniFrac_1.6	jpeg_0.1-9	rstatix_0.7.0
##	[82]	ggeffects_1.1.2	ggsignif_0.6.3	scales_1.2.0
##	[85]	memoise_2.0.1	plyr_1.8.7	zlibbioc_1.40.0
##	[88]	compiler_4.1.3	tinytex_0.39	clue_0.3-60
##		lme4_1.1-29	cli_3.3.0	ade4_1.7-19
##		XVector_0.34.0	htmlTable_2.4.1	Formula_1.2-4
##	[97]	MASS_7.3-57	mgcv_1.8-40	tidyselect_1.1.2
##	[100]	stringi_1.7.6	highr_0.9	yaml_2.3.5
##	[103]	locfit_1.5-9.5	latticeExtra_0.6-30	ggrepel_0.9.1
##		grid_4.1.3	tools_4.1.3	parallel_4.1.3
		rstudioapi_0.13	foreach_1.5.2	foreign_0.8-82
##	[112]	statip_0.2.3	gridExtra_2.3	$scatterplot3d_0.3-42$
##	[115]	farver_2.1.0	Rtsne_0.16	stable_1.1.6
##	[118]	digest_0.6.29	$shiny_1.7.1$	Rcpp_1.0.8.3
##	[121]	car_3.0-13	broom_0.8.0	performance_0.9.0
		later_1.2.0	httr_1.4.3	AnnotationDbi_1.56.1
		effectsize_0.7.0	sjstats_0.18.1	colorspace_2.0-3
		rvest_1.0.2	XML_3.99-0.10	fs_1.5.2
		modeest_2.4.0	truncnorm_1.0-8	splines_4.1.3
		yulab.utils_0.0.4	rmutil_1.1.9	$statmod_1.4.36$
		multtest_2.50.0	systemfonts_1.0.4	xtable_1.8-4
		jsonlite_1.8.0	nloptr_2.0.3	timeDate_3043.102
		R6_2.5.1	${\tt Hmisc\_4.7-1}$	NADA_1.6-1.1
		pillar_1.7.0	htmltools_0.5.2	mime_0.12
		glue_1.6.2	$fastmap_1.1.0$	$minqa_1.2.4$
		DT_0.23	BiocParallel_1.28.3	codetools_0.2-18
##	[157]	mvtnorm_1.1-3	utf8_1.2.2	lattice_0.20-45
		interp_1.1-3	survival_3.3-1	$biomformat_1.22.0$
		munsell_0.5.0	rhdf5_2.38.0	<pre>GenomeInfoDbData_1.2.7</pre>
		iterators_1.0.14	labelled_2.9.1	sjmisc_2.8.9
		haven_2.5.0	$reshape2_1.4.4$	gtable_0.3.0
##	[172]	bayestestR_0.12.1		