Abundance and short-term temporal variability of fecal microbiota in healthy dogs

Jose F. Garcia-Mazcorro¹, Scot E. Dowd², Jeffrey Poulsen¹, Jörg M. Steiner¹ & Jan S. Suchodolski¹

¹Gastrointestinal Laboratory, Department of Small Animal Clinical Sciences, College of Veterinary Medicine, Texas A&M University, College Station, TX
²Molecular Research LP (MR DNA), Shallowater, TX

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Abstract
Temporal variations of intestinal microorganisms have been investigated in humans, but limited information is available for other animal species. The aim of the study was to evaluate the abundance and short-term temporal variability of fecal microbiota in dogs. Two fecal samples were collected (15 days apart) from six healthy dogs. The microbiota was evaluated using fluorescence in situ hybridization (FISH) and 454-pyrosequencing targeting the 16S rRNA and its gene. Pyrosequencing revealed 15 families comprising >80% of all microbiota, over time intraindividual coefficients of variation (%CV) ranged from 2% to 141% (median: 55%). In contrast, the interindividual %CV ranged from 62% to 230% (median: 145%). Relative proportions of Faecalibacterium (important for intestinal health) and Subdoligranulum were low (two dogs harbored 4–7% of Subdoligranulum, the remaining dogs had <1% of either genus). Conversely, FISH revealed that Faecalibacterium comprised a median of 5% of total counts (range: 0–8%, probe Fprau645). A novel FISH probe (Faecali 698) was tested that, compared with Fprau645, can detect in silico a similar percentage of Faecalibacterium but higher proportions of Subdoligranulum. This probe revealed a high percentage of Faecalibacterium–Subdoligranulum (median: 16% of total counts). Future studies should consider the observed variability and discrepancies in microbial abundance between FISH and 454-pyrosequencing.

Introduction
The gastrointestinal (GI) tract of animals contains trillions of microorganisms that help maintain intestinal and overall health. Fueled by recent advances in the field of molecular tools and bioinformatics, the last decade witnessed an improved understanding on the phylogenetic structure of the intestinal microbiota and its symbiotic association with the host (Suchodolski 2011; Walter and Ley 2011).

The manipulation of intestinal microorganisms to improve health has been of interest for decades. However, the exact abundance and/or proportion of each member of the intestinal microbiota are a matter of debate, although some trends have been recognized for some bacterial groups (Hoyles and McCartney 2009). Also, only few studies have evaluated temporal variations of the intestinal microbiota (Franks et al. 1998; Buddington 2003; Matsuki et al. 2004; Dethlefsen and Relman 2011) and most of these studies have investigated the human microbiota only, with limited available information about the abundance and variability of the intestinal microbiota in other animal species. A recent study evaluated inter-individual differences in fecal microbiota of healthy cats and dogs using 454-pyrosequencing, but only one time point was evaluated and no truly quantitative assays were performed (Handl et al. 2011). Currently available microbial identification techniques can yield distinctive views of the membership composition of the intestinal microbiota, based on their differences in estimating the abundance of the microorganisms. For example, sequencing technologies, especially high-throughput techniques, such as microarray-based methods or 454-pyrosequencing, are able to characterize in depth the composition of the intestinal microbiota but are only semi-quantitative in nature. Other techniques, such as fluorescence in situ hybridization...
tion (FISH), can provide the actual numbers of the microorganisms but its application to all microbiota is technically challenging. The aim of this study was to evaluate the abundance and short-term temporal variability of fecal microbiota in a population of healthy dogs using a semi-quantitative (454-pyrosequencing) and a quantitative (FISH) approach.

**Materials and Methods**

**Animals and fecal collection**

Six privately owned pet dogs were enrolled (Table 1). All dogs were considered healthy on physical examination by a veterinarian, lived in different households, did not have any clinical signs of GI disease nor have consumed antibiotics for at least 3 months before fecal sample collection. Serum concentrations of cobalamin, folate, pancreatic lipase immunoreactivity, and trypsin-like immunoreactivity were measured to rule out subclinical GI disease (results not shown). Naturally passed fecal samples were collected from all dogs at two time points (15 days apart) within the same time frame (Fall 2010). The study protocol was approved by the Clinical Research Review Committee (CRRC#10-14) at Texas A&M University and written informed consent was obtained from the owners of all enrolled animals.

**Assessment of fecal microbiota**

**Extraction of DNA**

Genomic DNA was extracted and purified from 100 mg of each fecal sample using a phenol–chloroform extraction procedure as described by Suchodolski et al. (2005).

**454-Pyrosequencing**

The fecal bacterial taxonomic structure was evaluated using a bacterial tag-encoded FLX-titanium 16S rRNA gene amplicon pyrosequencing (bTEFAP) as described previously for canine fecal samples (Handl et al. 2011). The sequence processing pipeline was carried out as described in detail by Garcia-Mazcorro et al. (2012). Briefly, Phred20 quality reads were trimmed to remove tags and primer sequences, depleted of non-16S rRNA reads, chimeric, mitochondrial, and sequences of less than 250-bp length. The final sequences were evaluated using a standard nucleotide basic local alignment search tool (BLASTN) against a continually curated high-quality 16S rRNA gene database derived from the National Center for Biotechnology Information (NCBI). BLAST outputs were compiled to generate percentage files at each taxonomic level. Sequences with identity scores to known of well-characterized 16S rRNA gene sequences between 95% and 97% identity were resolved at the genus level, between 90% and 95% at the family level, between 85% and 90% at the order level, between 80% and 85% at the class level, and 77–80% at the phylum level. Sequence information is available through GenBank within a shortread archive (SRA) under accession (SRA053134).

**FISH**

For FISH analysis, we obtained an aliquot of 100 mg from each fecal sample and prepared paraffin-embedded fecal blocks (PEFB). These blocks were prepared because currently available FISH protocols often do not allow for a consistent and uniform distribution of the microorganisms across the glass slide and are associated with a low throughput. A detailed description of the procedure to prepare the fecal blocks from 100 mg of feces (wet weight) is provided as supplementary information.

For the FISH procedure, two serial sections (5 μm) of each fecal block were cut and placed on coated glass slides (IMEB Inc., San Marcos, California). The sections were deparaffinized by passage through xylene (3 × 10 min), 100% ethanol (2 × 5 min), 95% ethanol (1 × 5 min), and 75% ethanol (1 × 5 min). After the slides were air-dried, a total of 10 μL of warm hybridization buffer (containing 30 ng/μL of the respective probe, each labeled at the 5′ end with Cy5) was pipetted onto each section, covered by a cover glass, and incubated for 4 h at the optimum hybridization temperature (Table 2). The sections were washed with a wash buffer (hybridization buffer without sodium dodecyl sulfate [SDS]) for 30 min at 52°C. ProLong Gold antifade reagent (Invitrogen, Grand Island, New York) with 4',6-diamidino-2-phenylindole (DAPI, ~10 μL) was pipetted onto each section and sealed with transparent nail polish. All FISH probes used in this study are summarized in Table 2, which include the reverse complement of two recently developed polymerase chain reaction (PCR) sense primers to detect Faecalibacterium prausnitzii.

**Table 1.** Signalment of enrolled dogs.

| Dog | Breed          | Age   | Weight (kg) | Diet             |
|-----|----------------|-------|-------------|------------------|
| D1  | Labrador mix   | 10    | 33          | Beneful          |
| D2  | Boston Terrier | 4     | 11          | Science Diet     |
| D3  | Labrador mix   | 1.5   | 25          | Purina One       |
| D4  | Labrador mix   | 5     | 23          | Dry food         |
| D5  | Mixed          | 4     | 23          | Purina Sensitive |
| D6  | Australian Kelp | 9     | 16          | Purina High      |

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Table 2. Oligonucleotides used for FISH and PCR analyses.

| Sequence (5′–3′) | Target | Temperature (°C) | References |
|-----------------|--------|-----------------|------------|
| **FISH probes** |        |                 |            |
| Lab158          | GGTATAGCATCTGGTTCCA | Lactobacillus and Enterococcus spp. | 35 | Harmsen et al. (1999) |
| Erec482         | GCTTCCTAGTCAGTACCCG | Clostridium coccoideus–Eubacterium rectale group | 50 | Franks et al. (1998) |
| Fprau645        | CTCCTGCACTATCAAGAAAAAC | Faecalibacterium prausnitzii and relatives | 46 | Suau et al. (2001) |
| Faecali698      | GTGCCCGATGGCCGCTTCC | Faecalibacterium and relatives | 50 | This study |
| CFB555f         | CCCCCTAAAAACCAATDAAWTCGGG | Phylum Bacteroidetes | 50 | Muhling et al. (2008) |
| **PCR primers** |        |                 |            |
| Faecalif       | GAAGGCAGCTACTGGGCAC | Faecalibacterium and relatives | 60 | Garcia-Mazcorro et al. (2012) |
| FaecalIR       | GTGCAAGGCGAGTTGAGGCTG | Faecalibacterium spp. |        | |

1As described by listed references (published oligonucleotides) or the Ribosomal Database Project (RDP).
2Hybridization or annealing temperature.
3Based on the current RDP, this probe is specific only for Lactobacillus spp.
4This sequence is the reverse complement of the sense PCR primer (FaecaliF) and can detect in silico mainly Faecalibacterium spp. This oligonucleotide may also detect a proportion of phylogenetically related bacteria such as the genera Subdoligranulum (18% of all Subdoligranulum in RDP) and Anaerofilum (21% of all Anaerofilum in RDP). Anaerofilum has not been described in intestinal contents.
5This sequence is the reverse complement of the sense PCR primer to detect the phylum Bacteroidetes developed by Muhling et al. (2008).
6This sequence can only detect Faecalibacterium spp. in silico (based on RDP).

spp. (probe Faecali 698, see “Quantitative real-time PCR” below) and the Phylum Bacteroidetes (probe CFB555f; Muhling et al. 2008), respectively. We also evaluated the abundance of Faecalibacterium spp. (using the previously published probe Fprau645), Lactobacillus spp. (probe Lac 158), and the Clostridium coccoideus–Eubacterium rectale group (probe Erec 482) (Table 2). Images were obtained from a total of 20 random microscopic fields per time point analyzed (10 microscopic fields were obtained from each section of the fecal blocks) using a Zeiss Stallion digital confocal microscope (Carl Zeiss Microimaging Inc., Thornwood, New York) with a C-apochromat (63× water correction) objective lens. The fecal microorganisms were quantified using an image analysis software in the public domain (ImageJ: http://rsbweb.nih.gov/ij/index.html). One bacterial group (i.e., Lactobacillus spp., see below) was either absent or only showed low abundance in most dogs and was therefore counted manually instead. The following formula was used to calculate the number of bacterial cells per gram of wet weight of feces for each microscopic field:

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\text{Bacterial cells/g feces} = \text{bacterial cells in the microscopic field} \times 33,859 \times 600 \times 10,\quad\text{where 33,859 is the approximate number of microscopic fields in each paraffin slice of the fecal block (each microscopic field was 14,767 \mu m^2 and each paraffin slice is 500 million \mu m^2), 600 is the approximate number of paraffin slices (5 \mu m thick) in each fecal block (~3 mm deep), and 10 is the factor to multiply for to obtain the numbers of 1 gram of feces (each fecal block was made from 100 mg of feces). In an effort to reduce toxic waste, all hybridizations were performed without formamide in the hybridization buffer. Increasing concentrations of formamide were added to the hybridization buffer only to confirm the obtained counts with different stringency conditions and only in a subset of samples (see below). The probes Faecali 698 and CFB555f were tested for specificity against known strains of Faecalibacterium, Streptococcus, Enterococcus (phyllum Firmicutes), and Bifidobacterium spp. (phyllum Actinobacteria).

Quantitative real-time PCR

To verify the presence of Faecalibacterium in the fecal samples, bacterial DNA was amplified using primers for the 16S rRNA gene of Faecalibacterium spp. recently developed in our laboratory (Garcia-Mazcorro et al. 2012; Table 2). The PCR mastermix (10 \mu L) contained 5 \mu L of SsoFast EvaGreen supermix (Biorad Laboratories, Hercules, California), 2.2 \mu L of sterile water, 0.4 \mu L of each primer (final concentration: 200 nmol), and 2 \mu L of DNA (5 ng/\mu L). A melt-curve analysis was performed after all PCR cycles were terminated to confirm the specificity of the primers. The identity (Faecalibacterium spp.) of the amplicons was verified using an automated sequence analyzer (ABI PRISM 377 DNA Sequencer, Applied Biosystems, Foster City, California).

Statistical analysis

The coefficient of variation (%CV) was calculated using the following formula: [(standard deviation/average) × 100]. The standard deviation (squared root of variance) was calculated differently to obtain intra- and interindividual %CV. All FISH measurements for each
bacterial group were organized in spreadsheets with 12 columns (six dogs, two time points) and 20 rows (one for each microscopic field). Intraindividual %CV was calculated using standard deviations from each pair of FISH measurements within a dog (total of 20 pairs, one for each microscopic field). Interindividual %CV was calculated using standard deviations from each set of 12 FISH measurements (six dogs, two time points) within each row (total of 20 sets of 12 FISH measurements). Total %CV was calculated using all 240 FISH measurements (six dogs × two time points × 20 microscopic fields). Similar calculations were employed to obtain intra- and interindividual CV for the relative abundance of pyrosequencing tags (percentage of sequences) using data from the 15 most abundant bacterial families (≥80% of all sequences). The results are reported as medians with interquartile ranges.

Results

bTEFAP

The fecal microbiota of all enrolled dogs was dominated by organisms belonging to the phylum Firmicutes, which accounted for ≥75% of all sequences in all dogs (median: 88%, range: 75–98%) (Fig. 1). The second most abundant phylum was Actinobacteria (median: 3%, range: 1–22%), followed by Proteobacteria (median: 1%, range: 0–17%), Bacteroidetes (median: 1%, range: 0–7%), Fusobacteria (only three dogs harbored this phylum at one or both time points with <1% abundance), and Acidobacteria (only three dogs harbored this phylum at one or both time points with <0.1% abundance).

Fifteen families comprised ≥80% of the fecal bacterial microbiota in all dogs, and ≥90% of all microbiota in five of the six dogs (Fig. 2). From these 15 families, 10 (67%) families belonged to the phylum Firmicutes, two (13%) families to the phylum Bacteroidetes, two (13%) families to Actinobacteria, and one (7%) family to the phylum Proteobacteria (Fig. 2). The intraindividual %CV (over time) ranged from 2% to 141% (overall median: 55%). In contrast, the interindividual %CV ranged from 62% (family Ruminococcaceae) to 230% (family Lactobacillaceae) (overall median: 145%) (Table S1). Interestingly, the genus Lactobacillus was found to be highly abundant in one dog at both time points (Fig. 2, Dog 4, 73% and 75%, respectively, all other dogs had <1%), an observation that was not confirmed using FISH (see “FISH” results below). Clostridium was the most abundant genus in all dogs (median: 21%, range: 1–42%) and had an interindividual %CV of 84%. The genera Faecalibacterium and its phylogenetically related Subdoligranulum showed low abundance in most dogs (only two dogs had 4–7% of Subdoligranulum; the rest had ≤1% of either genus) and similarly variable among individual dogs (141 and 143 %CV, respectively). The genus Anaerofilum, a bacterial group that is phylogenetically related to both Faecalibacterium and Subdoligranulum, was detected only at one time point in one dog (<0.1% of all pyrosequencing reads).

FISH

All FISH results are summarized in Table 3. All dogs had detectable counts of all analyzed bacterial groups except for Lactobacillus spp. (probe Lac158), which was either absent (two dogs) or showed low abundance (three dogs harbored...
<1% of all microbiota at either time point). Only one dog showed relatively high counts of Lactobacillus (Dog 2 harbored \(5 \times 10^9\) counts at one time point, or about 2% of all microbiota, while at the other time point <1% of total counts were observed). These results are not in agreement with the pyrosequencing data (see above), where all dogs had <1% of Lactobacillus except for Dog 4, who harbored >70% of Lactobacillus at both time points.

All dogs showed detectable counts of Faecalibacterium spp. using both FISH probes with the exception of Dog 3, which did not have any Faecalibacterium spp. as estimated using the FISH probe Fprau645. This dog also did not harbor detectable Faecalibacterium sequences using quantitative real-time PCR (see below). The use of the probe Fprau645 revealed an overall median of Faecalibacterium of 5% of DAPI counts (range: 2–7%), whereas the novel probe Faecali 698 yielded about three times those counts (median: 16% of DAPI counts, range: 10–22%) (Table 3). This difference in proportions between the two FISH probes may be due to populations of Subdoligranulum and Anaerofilum, although Anaerofilum has not been described in intestinal contents (see “Discussion” below). The intra- and interindividual variability was similar for both probes (Table 3).

The numbers and proportions of Faecalibacterium spp. (probe Faecali 698) and the phylum Bacteroidetes (probe CFB555f) for subjects harboring the highest amounts of these bacteria were confirmed using increasing concentrations of formamide (results not shown). The specificity of these probes was verified in vitro against strains of Lactobacillus, Bifidobacterium, Streptococcus, and Enterococcus spp. All bacteria were fluorescent when using a universal probe (Eub 338) and did not fluoresce when using these two probes.

### Quantitative real-time PCR

Quantitative real-time PCR was used to verify the presence of Faecalibacterium in the fecal samples. With the exception of one dog (Dog 3), the use of our Faecalibacterium-specific primers yielded the expected 598-bp amplicon in all the samples analyzed. This amplicon corresponds to the positions 698–1291 of the 16S rRNA gene sequence of Faecalibacterium prausnitzii (ATCC 27766).

### Discussion

Temporal variations of the intestinal microbiota have been investigated in humans, but limited information is available for other animal species. Also, different microbial identification techniques can yield differing results of the composition of the intestinal microbiota, based on their differences in estimating the abundance of the microorganisms. This study investigated the abundance and short-term temporal variability of fecal microbiota using pyrosequencing and FISH in a population of healthy dogs.

Pyrosequencing (as evaluated using bTEFAP) revealed a distinctive relative abundance of the fecal microbiota in each of the enrolled dogs (Fig. 1), especially at the family level (Fig. 2). As suggested by others (Matsuki et al. 2004; Suchodolski et al. 2008), our results support the possibility that the variation of the intestinal microbiota among individuals (overall median %CV: 145% in this study) is higher than the variation within individuals over time (overall median %CV: 55% in this study). These observations are likely due to the fact that individual dogs had little in common and were exposed to the same environmental conditions (within dogs) during the 15 days period of fecal sample collection. Interestingly, the family with the lowest interindividual variability in abundance (i.e., Ruminococcaceae) has been proposed to be a member of the core human fecal microbiota (Tap et al. 2009), suggesting that some intestinal bacterial groups may also be consistently shared not only among humans but also among other mammals. However, the presence of a core

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**Table 3. Summary of quantitative results obtained from the FISH analyses.**

| FISH probe | Abundance of the bacteria (median with interquartile range) | Percentage of DAPI counts | %CV (median with interquartile range) |
|------------|-----------------------------------------------------------|---------------------------|-------------------------------------|
|            | Number of cells/g wet feces Log_{10} cells/g wet feces | Intraindividual Interindividual Total %CV |                                    |
| Erec482    | 3.7 (2.2–4.9) \( \times 10^9\) 9.6 (9.3–9.7) 22 (13–27) | 40 (17–59) 48 (43–54) 48% |                                    |
| CFB555f    | 4.7 (2.8–9.5) \( \times 10^9\) 9.7 (9.5–10.0) 28 (15–41) | 38 (18–64) 75 (67–84) 77% |                                    |
| Fprau645   | 1.0 (4.1 \( \times 10^9\)–1.6 \( \times 10^9\)) \( \times 10^9\) 9.0 (8.8–9.2) 5 (2–7) | 28 (11–71) 99 (87–106) 96% |                                    |
| Faecali 698| 2.8 (1.7–5.1) \( \times 10^9\) 9.5 (9.2–9.7) 16 (10–22) | 28 (13–56) 68 (65–79) 70% |                                    |

1The genus Lactobacillus was also quantified using FISH but showed very low abundance (<=2%) in all dogs. Therefore, it is not included here.

2The median and interquartile ranges of the abundance of each bacterial group were calculated from a total of 240 observations (six dogs, two time points, 20 microscopic fields each).

3The median and interquartile ranges of the coefficients of variation (%CV) were calculated from measurements between the two time points (intraindividual) and among individuals (interindividual). Total %CV was calculated from all 240 observations.
fecal microbiota in nonhuman individuals has not been investigated to date.

The probe EreC482 has previously been used to evaluate the abundance of the Clostridium coccoidei–Eubacterium rectale (Erec) group in several human studies (Marteau et al. 2001; Matsuki et al. 2004; Sokol et al. 2006). Also, one recent study showed that dogs may harbor 9.2–9.6 log_{10} cells/g of wet feces (~10% of all fecal microbiota, as estimated by DAPI staining) of the Erec group (Jia et al. 2010). Similarly, this study showed an overall median of 3.7 × 10^8 (9.6 log_{10}) cells per g of wet feces of the Erec group (overall median: 22% of all fecal microbiota as estimated using DAPI staining). The difference in the proportions of the Erec group between the study by Jia et al. (10%) and this study (22%) may be due to interindividual differences in total fecal bacterial counts as well as to differences in the study populations. For example, Jia et al. used only Beagle dogs, whereas this study used a more heterogeneous population of dogs. Because the Erec group consists of members of different bacterial groups within the Firmicutes, it is difficult to try to estimate its true proportion based on pyrosequencing results. For example, an in silico analysis of specificity using the Probe-Match tool of RDP (all analyses using probe match were performed with 0 errors allowed) shows that the EreC 482 probe can only detect the genus Blautia (family Incertae sedis XIV) and several genera within the family Lachnospiraceae (e.g., Butyrivibrio, Coprococcus, Roseburia, Dorea, Anaerostipes). Together, all these genera comprise ~4% of all pyrosequencing tags in this study. This matter is further complicated because of recent reclassifications of some of these bacterial groups (e.g., Liu et al. 2008).

Recently, the phylum Bacteroidetes has received considerable interest due to its potential involvement in the pathogenesis of obesity (Ley et al. 2005; Turnbaugh et al. 2006). One recent study showed that this phylum may be more abundant (~2% of all fecal microbiota) in the feces of dogs (Jia et al. 2010). Similarly, this study showed an overall median of 3.7 × 10^8 (9.6 log_{10}) cells per g of wet feces of the Bacteroidetes group (overall median: 14% of all fecal microbiota as estimated using DAPI staining). Published results in humans with respect to the proportion of the Bacteroidetes also vary widely (Armougom and Raoult 2008). Studies are needed to determine the identities and exact proportions of members from the Bacteroidetes in feces of dogs.

The bacterium Faecalibacterium prausnitzii has received increased attention because of its potential protective role against inflammatory bowel disease in humans (Sokol et al. 2008). Interestingly, one study suggested that the Faecalibacterium species found in dogs may not be F. prausnitzii (Suchodolski et al. 2008), based on the phylogenetic affiliation of near-full-length 16S rRNA gene sequences belonging to a canine clone (C2-02) and a human strain (AJ270469). We have designed two PCR oligonucleotides that are capable of amplifying the genus Faecalibacterium (as evaluated by in silico analysis and sequencing of amplicons, Table 2). From these two primers, we chose the sense primer (FaecalilF) as candidate for a FISH probe because it can in silico detect 22% more Faecalibacterium sequences than the anti-sense primer (81% vs. 59% of all Faecalibacterium sequences in RDP). However, this sense primer may also detect some phylogenetically related bacteria, the genus Subdoligranulum (Holmstrom et al. 2004) and the poorly described genus Anaerofilum (Zellner et al. 1996). Importantly, the presence of Anaerofilum in intestinal contents has not been described in the literature. To our knowledge, the only other FISH probe available to detect intestinal Faecalibacterium is the probe Fprau645 that was developed by Suau et al. (2001). This probe does not detect any Subdoligranulum or Anaerofilum (based on RDP) and can detect in silico many Faecalibacterium (78% of all Faecalibacterium sequences in RDP). Using this probe, Suau et al. (2001) suggested that F. prausnitzii and its relatives may comprise ~17% of all fecal microbiota (as estimated by DAPI staining) of healthy humans. Using the same FISH probe, another, more recent study, also showed that Faecalibacterium and relatives comprise 14–17% of all fecal microbiota in humans (Benus et al. 2010). In contrast, in this study, the use of this FISH probe revealed that Faecalibacterium and relatives comprise ~5% of all fecal microbiota in the enrolled dogs, while our FISH probe revealed about three times this proportion (~16% of all microbiota). This difference in proportions between the two FISH probes may be due to populations of Subdoligranulum. Interestingly, pyrosequencing revealed that both Faecalibacterium and Subdoligranulum were present in low abundance in most dogs (only two dogs had 4–7% of Subdoligranulum, the rest of
the dogs had ≤1% of either genus), whereas the genus *Anaerofilum* was detected only in one dog and at one time point (<0.1% of all pyrosequencing reads). As with other bacterial groups, studies are needed to determine the exact abundance of Faecalibacterium and its relatives to help determine their role during intestinal disease in dogs and other animal species.

This study shows some discrepancies between the pyrosequencing and the FISH results, especially with regard to the abundance of the bacterial groups. This was expected, in part, because pyrosequencing relies on PCR to amplify genomic targets and may therefore underestimate the true abundance of all members of the microbiota. Also, the primers used in this study for pyrosequencing can amplify the variable regions V1 to V3 of the 16S rRNA gene, whereas some of our FISH probes (e.g., Faecali 698) target regions outside of these regions. However, 454-pyrosequencing and other high-throughput techniques can provide the relative proportions of the different bacterial groups, which ultimately provide useful information with regard to a change in the relative abundance of the microorganisms.

This study used two oligonucleotides to determine the abundance of the phylum Bacteroidetes (probe CFB555f) and Faecalibacterium relatives (probe Faecal 698) in feces of dogs using FISH. However, the ability of these two probes to hybridize with related and unrelated bacteria has been only evaluated *in silico* either by our group (Faecal 698, this study) and by Muhling et al. (2008) (probe CFB555f). Muhling et al. went a step further and created a clone library using a primer pair containing this oligonucleotide, and showed that all sequences produced in the clone libraries were derived from members of the Bacteroidetes. In this study, we additionally showed that these two probes do not hybridize with members of two different phyla *in vitro*, the Firmicutes (*Lactobacillus*, *Streptococcus*, and *Enterococcus* spp.) and the Actinobacteria (*Bifidobacterium* spp.). Further studies are nevertheless warranted to investigate more extensively the specificity *in vitro* of these two oligonucleotides using FISH.

In summary, normal temporal variations of the intestinal microbiota have been studied mainly in humans, with little information available in other animal species. Pyrosequencing and FISH are different molecular techniques that are commonly used in gut microbial ecology to characterize the intestinal microbiota and its changes in response to external influences. Our results suggest that these two techniques yield different abundances of some of the most predominant fecal microbiota in healthy dogs, including Faecalibacterium, a bacterial group that has consistently been shown to be depleted during intestinal disease, and Bacteroidetes, a phylum which has been suggested to play an important role in energy harvesting and obesity. These abundances seem to vary more among dogs than within dogs within a period of 2 weeks, although this seems also to be dependent on the technique employed and the specific bacterial group analyzed. These observations should be considered when evaluating the effect of agents on the canine intestinal microbiota.

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**Conflict of Interest**

None declared.

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**Supporting Information**

Additional Supporting Information may be found in the online version of this article:

**Appendix S1.** Instructions to prepare fecal blocks from 100 mg of feces.

**Table S1.** Relative abundance (proportions of pyrosequencing reads) of the most predominant bacterial families in fecal samples from six dogs (D1–D6) and at two time points 15 days apart (-1 and -2) with interindividual coefficients of variation (%CV).

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