Evaluating DNA Extraction Methods for Community Profiling of Pig Hindgut Microbial Community

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Abstract

Recovery of high quality PCR-amplifiable DNA has been the general minimal requirement for DNA extraction methods for bulk molecular analysis. However, modern high throughput community profiling technologies are more sensitive to representativeness and reproducibility of DNA extraction method. Here, we assess the impact of three DNA extraction methods (with different levels of extraction harshness) for assessing hindgut microbiomes from pigs fed with different diets (with different physical properties). DNA extraction from each sample was performed in three technical replicates for each extraction method and sequenced by 16S rRNA amplicon sequencing. Host was the primary driver of molecular sequencing outcomes, particularly on samples analysed by wheat based diets, but higher variability, with one failed extraction occurred on samples from a barley fed pig. Based on these results, an effective method will enable reproducible and quality outcomes on a range of samples, whereas an ineffective method will fail to generate extract, but host (rather than extraction method) remains the primary factor.

Introduction

The vertebrate hindgut microbiome is critical to host organism nutrition, health, and welfare, including control of infectious disease [1–2]. Faecal samples are used as a simple, non-invasive method of sampling this community [3]. While the hindgut microbiome has been extensively assessed in humans [4], it is also highly important in other animals, including domestic and commercial livestock [5]. This is not only for commercial and welfare reasons, but because large animals such as pigs are being increasingly used as models for human microbiomes [6]. The most common culture-independent method for analysing microbial communities is 16S rRNA amplicon profiling [7] which can be affected by the DNA extraction method [8]. Several studies have attempted to develop or validate DNA extraction methods suitable for faecal samples. Clement et al. [9] modified the UltraClean Soil DNA kit (Mo Bio Laboratories, Solana Beach, CA, USA) with dry lysis tubes and a second DNA wash step to produce a high yield of
PCR-quality DNA from human faeces. Tang et al. [10] also described a modified method, utilizing hexadecyltrimethylammonium bromide (CTAB), salt, polyvinylpyrrolidone and betamercaptoethanol for cell lysis and chloroform for DNA isolation, producing notably better results than the QIAamp DNA stool mini kit. Salonen et al. [11] also concluded that a DNA extraction method using repeated bead beating [12] can generate up to 35-fold increase on DNA yield than other extraction method with non or less mechanical cell lysis.

DNA yield is most commonly used as a proxy for method quality, the assumption being that a higher yield is more representative of the community under study. However, the severity of extraction is an important factor affecting the representativeness and reproducibility of extraction methods. Overly harsh methods while generally producing higher yields can potentially degrade the DNA of sensitive organisms (e.g. Bacteroidetes, [13]), while excessively gentle methods can fail to extract from gram-positive organisms with thick cell walls (e.g. Clostridium, [11]). Reproducibility of extraction can also be an issue, particularly in livestock faecal samples, which can have a high degree of heterogeneity due to the faecal matrix, particularly where diet is varied.

The development of next-generation sequencing (NGS) provides a powerful tool to increase depth and resolution of community analysis [7] and NGS has been applied for microbial profiling of faecal sample from humans [14–16] or animals such as swine [5], white rhinoceros [17] and horses [18]. Due to its enhanced depth and resolution, NGS techniques can be more influenced or likely to detect artefacts due to DNA extraction methods, and hence analysis of the impacts of extraction is an important consideration. Recently, the variations resulting from choice of extraction method were shown to be significant in NGS study on human gut [13] but negligible on insects gut [19], while this remains unclear on livestock gut.

Here, we evaluate three commonly used pig faecal DNA extraction methods (FastDNA SPIN kit, PowerSoil kit and a previous reported protocol utilizing CTAB) using NGS sequencing. All three methods gave comparable results despite striking differences in DNA yields.

Materials and Methods

Pig Faecal samples

Fresh whole faecal samples were collected from three pigs aged 11 weeks, as part of a larger metabolic trial being conducted at the Advanced Animal Science facility of the University of Queensland (Gatton, Queensland, Australia) with permission from the University of Queensland. Animal ethics approval for the larger metabolic trial (from which faecal samples were obtained) was provided by the Staff Access Animal Ethics Committee, Argri-Science Queensland. Sampling procedures were reviewed and specifically approved as part of the approval.

Faecal samples were collected in sterile bags and stored on ice, immediately taken to the laboratory and subsequently frozen at -20°C. No invasive sampling or sacrificing was involved in sample collection. Pig diets were the three most common commercial Australian carbohydrate based diets and are provided in supplementary material (S1 Table) with mainly wheat fines in pig A (72%) and pig B (65%). The diet for pig C was replaced with barley fines (48%).

DNA extraction

Samples were homogenized to mix the liquid, mud layers and solid settlement and subsampled. DNA was extracted from 0.3 g aliquots in triplicate using two commercially-available kits: FastDNA SPIN Kit for Soil (MP Biomedicals, Santa Ana, California, US) referred as FAS, PowerSoil® DNA Isolation Kit (MoBio Laboratories, Carlsbad, CA, US) referred as POW and one additional protocol utilizing CTAB adapted from Tang et al. [10] referred as CON. The
concentration of each eluted DNA was measured by Nanodrop spectrometer (Thermo Scientific, US) as well as the purity (indicated as 260/280 ratio). The extracted DNA was loaded on a 1% agarose gel to identify extent of DNA degradation.

**FAS method.** DNA extraction with FAS was performed according to manufacturer’s instruction with modification on the bead beating time and additional pre-elution incubation (details below). Sample aliquot was added into the lysing matrix with lysis buffer supplied. Lysing matrix was blended with mini Bead Beater (BioSpec, Bartlesville, US) at 4,800 oscillations per minute for 60 seconds. After being cooled to 4°C, tubes were centrifuged at 13,000 x g for 15 minutes to pellet debris. Supernatant was transferred to 250 µL protein precipitation solution and mixed with 1 mL binding matrix. The mixture of binding matrix and DNA was then filtered and washed with 500 µL SEWS-M supplied in the kit. Additional incubation at 50°C for 5 minutes was performed before the final elution. 50 µL RNAnase-free water was used to elute the DNA from filter.

**POW method.** DNA extraction with POW was performed according to the manufacturer’s instructions with modifications of bead beating time and additional pre-elution incubation. Sample aliquot was added into the lysing matrix with lysis buffer supplied. Lysing matrix was then blended with mini Bead Beater (BioSpec, Bartlesville, US) at 4,800 oscillations per minute for 60 seconds. The remaining steps (including protein removal and DNA washing) were performed as recommended. DNA was also eluted with 50 µL RNAnase-free water with pre-elution incubation at 50°C for 5 minutes.

**CON method.** The conventional DNA extraction method used in this study was described by Tang et al. (2008), in which autoclaved beads (0.5 g, 0.3 mm in diameter) were mixed with 570 µL buffer TE in capped tubes. Sample aliquot was then added. After bead beating at 4,800 oscillations per minute for 60 seconds, 5 µL 10% SDS and 3 µL Proteinase K was added to the tube. Tubes were incubated at 37°C for 1 hour and then at 65°C for 10 minutes. Supernatant (600 µL) was transferred to a clean autoclaved 1.5 mL tube with 100 µL 5M NaCl and 80 uL CTAB (65°C). Following this, Phenol:Chlorophom: Isoamyl-alcohol (800 uL, 25:24:2) were added to the tube and centrifuged at 13,000 x g for 5 minutes. The aqueous phase was then transferred to a new autoclaved Eppendorf tube. An equal volume of chloroform/Isoamyl-alcohol (50/50 v/v) was then added and mixed. After centrifuging at 13,000 x g for 5 minutes, the aqueous phase was then transferred to another autoclaved tube with 300 µL isopropanol. This was incubated at -20°C for 30 minutes. After centrifuging at 4°C, 13,000 x g for half an hour, the supernatant was removed and 500 µL 70% ethanol was added. The tube was again centrifuged at 4°C, 13,000 x g for 5 minutes. The supernatant was then removed. After the pellet was air dried, DNA pellet was resuspended in 50 µL RNAnase-free water.

**Pyrosequencing and data analysis**

DNA samples (300 ng each) were provided to Australian Centre for Ecogenomics (ACE) for pyrosequencing analysis. The primers used for pyrosequencing were 926f (5’ – AAACCTACAGGCTATGATATGC-3’) [20] and 1392r (5′ –ACGGGCGGTGTGTAC-3′) [21] targeting V6-V8 regions of the SSU rRNA gene with Roche 454 GS FLX sequencer (Roche, Switzerland).

Pyrosequencing results were analysed through the ACE Pyrosequencing Pipeline (https://github.com/minillinim/APP). Sequences reads were split according to barcodes in QIIME v1.8.0 [22]. De-multiplexed sequences were then trimmed to 250bp and de-noised by ACACIA [23]. Sequences with ≥97% similarity were assigned to operational taxonomic units (OTUs) by CD-HIT-OTU [24–25] and aligned by Pynast [26]. Each sequence was then classified using BlastTaxonAssigner in QIIME against the Greengenes database (2012 October release). Weighted UniFrac distances [27] were also calculated in QIIME.
Non-normalized OTU tables and rarefaction curves were generated by QIIME. An in-house
script Normaliser (https://github.com/minillinim/Normaliser) was used to find a centroid nor-
malized OTU table. The normalized OTU table was imported into R (version 3.0.1, [28]) to
generate multidimensional scaling analysis using Bray-Curtis dissimilarity method with func-
tion metaMDS in Package “vegan” [29]. Analysis of variance (ANOVA) was performed on the
OTU table with function aov in package “stats” [28]. Power analysis of ANOVA was performed
with function pwr.anova.test in package “pwr” in R [30] with number of groups as 3, and
observations per group of 9. Eta-squared effect size was calculated according to [31] as: sum of
squares between groups / total sum of squares. Analysis of similarity (ANOSIM) was per-
formed in QIIME [22]. De-multiplexed pyrosequencing data were deposited in GenBank
under BioProject PRJNA286807.

Results and Discussion
DNA quantity and purity
DNA of faecal samples collected from pig A and B fed with mainly wheat fines and pig C fed
with mainly barley, were extracted using two commercial kits (FAS and POW) and one con-
tventional method (CON). With the exception of pig C extracted by CON, all other samples
recovered high purity DNA template with $A_{260}/A_{280}$ ratios being close to 1.8 (Fig 1). The
amount of DNA extracted from FAS was the highest with up to two and ten times the

Fig 1. Yield (bars), purity (●) and PCR amplifiability of extracted DNA from pigs A, B, and C by different extraction methods FAS, POW and CON.
FAS method yields more pure and amplifiable DNA than other methods. Yield (bars) is plotted on the left y-axis, purity (●) is plotted on the right y-axis. Error
bars represent standard deviation. PCR amplifiabilities are indicated as “✓” for successful amplification or “×” for failed amplification.
doi:10.1371/journal.pone.0142720.g001
concentration of DNA compared to CON and POW respectively. There was high variability in concentration of DNA extracted from CON in both pig B and C replicates as shown by a large error bar in Fig 1. POW gave consistently lowest but reproducible DNA concentration (10–20 ng μL⁻¹) in all samples. FAS resulted in discrete high molecular weight bands from pig A and B, but with degraded DNA from pig C (S1 Fig). Clear single bands were produced by POW on triplicates from all three samples. The lowest quality DNA was generated by CON with conspicuous degradation in both pig B and C.

All samples were successfully PCR-amplified with the exception of pig C extracted by CON, possibly due to protein or phenol contamination (A₂₆₀/A₂₈₀: 1.4), which can inhibit PCR [32]. CON has been previously described as an effective DNA isolation method from healthy multiparous Rongchang sow faeces [10] compared to QIAamp DNA Stool Mini Kit (QIAGEN, Duesseldorf, Germany) and three other conventional methods [33–35]. However, results show variation both in technical replicates, and across extraction methods. DNA yield, quality and success of PCR amplification have been previously used as DNA extraction method criteria [36–37]. Based on the combination of these three criteria, it would appear that FAS is the most consistent method to use for extracting DNA from porcine faeces (Fig 1).

Pyrosequencing result comparison

Pyrosequencing generated 132,984 high quality reads after quality filtering, which were grouped into 1,065 operational taxonomic units (OTUs). FAS resulted in lower numbers of OTUs from the same sample, especially from pig C (191±11, standard error), numbers of OTUs from other samples are listed in S2 Table. Nonmetric Multi-Dimensional Scaling (NMDS) analysis of the sequence data (weighted UniFrac distances) showed primary separation of microbial communities according to ANOVA by host (R² = 0.94, p = 5.2×10⁻¹², with a power of 0.98) and secondarily by DNA extraction method (p = 7.6×10⁻⁰³) (Fig 2A). Separation within factors as analysed by ANOSIM was also strong with separation factors for host (R = 0.7, p = 0.001), and DNA extraction method (R = 0.2, p = 0.005).

Differences between replicates from the same extraction method were generally smaller than the differences between different extraction methods (ANOSIM: R = 0.5, p = 0.03 for pig A; R = 0.5, p = 0.02 for pig B) (Fig 2B and 2C). The same conclusion was found by evaluating different extraction methods on mock and bronchoalveolar lavage samples [38]. FAS had the highest variability in technical replicates in pig A and B fed with mainly wheat fines (Fig 3A and 3B). Although the UniFrac distances between replicates extracted by FAS from pig C was the lowest compared to POW and differences between methods (Fig 3C), it produced smeared gels (S1 Fig) and recovered the least OTUs indicating partial destruction of the DNA. Reproducibility from CON was also not stable as indicated by a large interquartile range (IQR, Fig 3B). CON utilized CTAB as the major lysis reagent in addition to bead beating and lysozyme (which were also used in both POW and FAS), which has been previously shown to generate variability in technical replicates [11]. In general, POW resulted in a lower range within replicates, and with a lower IQR, but this was not consistent for all hosts. In addition, pig C had the highest UniFrac distance and IQR indicating large variation between DNA extracted from FAS and POW. The major components in diet fed to pig C are barley fines which are physically more granular and abrasive than wheat fines [39], and thus possibly contribute to additional physical shear in addition to the extraction method.

Specific OTUs in the profiled communities can be identified which represent easier (Gram-negative) and harder (Gram-positive) to lyse populations (Fig 4). Consistent with NMDS analysis of microbial profiles from individual hosts (Fig 2B and 2C), POW and CON produced similar OTU relative abundances with FAS being the outlier method, increasing relative
abundance of Gram-positive organisms and decreasing that of Gram-negatives (Fig 4). Abundance of Gram-positive bacteria recovered from FAS were often higher than other methods, which is likely due to two factors: 1. Because of thicker cell walls, and large amounts of peptidoglycan, Gram-positive bacteria are difficult to physically lyse. FAS is a harsher method and hence able to recover more Gram-positive microbes. 2. The abundance of Gram-negative bacteria such as *Prevotella* (of pig C in Fig 4) is relatively lower due to DNA destruction and hence leads to a relative increase in Gram-positive bacteria such as *Clostridium*, *Streptococcus*, and *Lactobacillus*. Similar effects were identified in other pigs with low abundance (S3 Table). The latter is likely, as FAS recovered the least OTUs and produced visible smeared gels from pig C. The same reduction on *Escherichia* (Gram-negative bacterium) can be observed to a lesser extent, except from pig A due to an outlier replicate with extreme high abundance (13% compared to 3 and 4% in other replicates of pig A, S3 Table), indicating a high variability between replication by FAS. Therefore, although FAS appeared to be the best method in terms of yield purity and PCR amplifiability (Fig 1), community profiling data suggest that POW produces less bias and should be the preferred method. A harsher single extraction such as FAS may be preferable for ensuring hard to lyse populations are accessed, but less suitable for objective community assessment. While CON was the only non-commercial method, it was the least

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Fig 2. Nonmetric multidimensional scaling analysis of pyrosequencing result extracted with different methods between different pigs (A) within pig A (B) and within pig B (C). The replicates of each pig clustered well. Pyrosequencing results obtained from FAS differed to other methods in all pigs. Ellipses represent 95% prediction intervals (s_d.025_n-1), including correlation for host (A) or methods (B & C).

doi:10.1371/journal.pone.0142720.g002
suitable in general for these samples, and the cost of commercial kits is relatively minor in comparison with sampling and sequencing costs.

Conclusions

The main factor determining microbial community profile variation is host, rather than DNA extraction method, indicating that the tested extraction methods likely do not substantially skew community composition. Similar conclusions were reported in human gut microbiota profiling [40]. Samples from which extraction is difficult (because of the matrix, i.e. pig C fed on barley) can be extracted using a harsher method, but variability in replicates, and between
extraction methods increases, and can possibly lead to overestimation of the relative abundance of Gram-positive bacteria. The PowerSoil™ DNA Isolation Kit (POW), which extracts moderate yields of high quality DNA without apparently over-representing Gram-positive OTUs, is recommended for community profiling of pig faecal samples.

**Supporting Information**

S1 Fig. Example of DNA quality from different methods (FAS, POW and CON) and pigs (A, B and C) as shown in 2D-gel electrophoresis.

S1 Table. Proportions of feed ingredients for pigs.

S2 Table. Number of OTUs recovered from each sample.

S3 Table. Relative abundance of OTUs in the replicates of pigs.

**Acknowledgments**

We are grateful to Alan Skerman for providing samples to this study and Adam Skarszewski for the assistance on ACE Pyrosequencing Pipeline usage. We also thank Fiona May for performing the 454 pyrosequencing.

**Author Contributions**

Conceived and designed the experiments: YL PH DB. Performed the experiments: YL. Analyzed the data: YL DB. Contributed reagents/materials/analysis tools: PH DB. Wrote the paper: YL.
References

1. Muegge BD, Kuczynski J, Knights D, Clemente JC, González A, Fontana L, et al. Diet drives convergence in gut microbiome functions across mammalian phylogeny and within humans. Science. 2011; 332(6032):970–974. doi:10.1126/science.1198719 PMID: 21596990

2. De Jong A, Smee A, Ludwig C, Stephan B, De Graef E, Vanrobaeys M, et al. Antimicrobial susceptibility of Salmonella isolates from healthy pigs and chickens. Vet Microbiol. 2014; 171(3–4):289–306. doi:10.1016/j.vetmic.2014.01.030 PMID: 24598135

3. Beja-pereira A, Oliveira R, Alves PC, Schwartz MK, Luikart G. Advancing ecological understandings through technological transformations in noninvasive genetics. Mol Ecol Resour. 2009; 9(5):1279–1301. doi:10.1111/j.1755-0998.2009.02699.x PMID: 21564900

4. Huttenhower C, Gevers D, Knight R, Abubucker S, Badger JH, Chinwalla AT, et al. Structure, function and diversity of the healthy human microbiome. Nature. 2012; 486:207–214. doi:10.1038/nature11234 PMID: 22699609

5. Lamendella R, Domingo JWS, Ghosh S, Martinson J, Oerther DB. Comparative fecal metagenomics unveils unique functional capacity of the swine gut. BMC Microbiology. 2011; 11:103. doi:10.1186/1471-2180-11-103 PMID: 21575148

6. Zhang Q, Widmer G, Tzipori S. A pig model of the human gastrointestinal tract. Gut Microbes. 2013; 4(3):193–200. doi:10.4161/gmic.23967 PMID: 23549377

7. Tringe SG, Hugenholtz P. A renaissance for the pioneering 16 S rRNA gene. Curr Opin Microbiol. 2008; 11(5): 442–446. doi:10.1016/j.mib.2008.09.011 PMID: 18817891

8. Morgan JL, Darling AE, Eisen JA. Metagenomic sequencing of an In vitro-simulated microbial community. PLoS ONE. 2010; 5(4):e10209. doi:10.1371/journal.pone.0010209 PMID: 20419134

9. Clement BG, Kitts CL. Isolating PCR-quality DNA from human feces with a soil DNA kit. Biotechnology. 2000; 28(4):640–646.

10. Tang J, Zeng Z, Wang H, Yang T, Zhang P, Li YL, et al. An effective method for isolation of DNA from pig faeces and comparison of five different methods. J Microbial Methods. 2008; 75(3):432–436. doi:10.1016/j.mimet.2008.07.014 PMID: 18700153

11. Hair D, Frear J, Moss TM, Henrichs S, Nel T, de Kock DJ, et al. Microbiota and metabolic effects fromponse of the human gut to dietary intervention: a randomised controlled trial. Gut Microbes. 2015; 6(1):110–120. doi:10.4161/gmic.34203 PMID: 25483160

12. Yu Z, Morrison M. Improved extraction of PCR-quality community DNA from digesta and fecal samples. Biotechniques. 2004; 36(5): 808–812. PMID: 15152600

13. Wesolowska-Andersen A, Bahl MI, Carvalho V, Kristiansen K, Sicheritz-ponten T, Gupta R, et al. Choice of bacterial DNA extraction method from fecal material influences community structure as evaluated by metagenomic analysis. Microbiome. 2014; 2:19. doi:10.1186/2049-2618-2-19 PMID: 24949196

14. Hooda S, Vester Boler BM, Rossoni Serao MC, Brulc JM, Staeger MA, Boileau TW, et al. 454 pyrosequencing reveals a shift in fecal microbiota of healthy adult men consuming polydextrose or soluble corn fiber. J Nutr. 2012; 142(7): 1259–1265. doi:10.3945/jn.112.158766 PMID: 22649263

15. Qin J, Li Y, Cai Z, Li S, Zhu J, Zheng F, et al. A metagenome-wide association study of gut microbiota in type 2 diabetes. Nature. 2012; 490(7418):55–60. doi:10.1038/nature11450 PMID: 23023125

16. Finegold SY, Dowd SE, Gontcharova V, Liu C, Henley KE, Wollcott RD, et al. Pyrosequencing study of fecal microflora of autistic and control children. Anaerobe. 2010; 16(4): 444–453. doi:10.1016/j.anaerobe.2010.06.008 PMID: 20603222

17. Bian G, Ma L, Su Y, Zhu W. The microbial community in the feces of the white Rhinoceros (Ceratotherium simum) as determined by barcoded pyrosequencing analysis. PLoS ONE. 2013; 8(7): e70103. doi:10.1371/journal.pone.0070103 PMID: 23922920

18. Dougal K, de la Fuente G, Harris PA, Girdwood SE, Pinloche E, Geor RJ, et al. Characterisation of the faecal bacterial community in adult and elderly horses fed a high fibre, high oil or high starch diet using 454 Pyrosequencing. PLoS ONE. 2014; 9(2): e87424. doi:10.1371/journal.pone.0087424 PMID: 24504261

19. Rubin BER, Sanders JG, Hampton-marcell J, Owens SM, Gilbert JA, Moreau CS. DNA extraction protocols cause differences in 16S rRNA amplicon sequencing efficiency but not in community profile composition or structure. MicrobiologyOpen. 2014; 3(6):910–921. doi:10.1002/mbo3.216 PMID: 25275543

20. Lane DJ. 16S/23S rRNA sequencing. In: Stackebrandt E, Goodfellow M, editors. Nucleic acid techniques in bacterial systematics. John Wiley and Sons: New York; 1991. pp 115–175.
21. Lane DJ, Pace B, Olsen GJ, Stahl DA, Sogin ML, Pace NR. Rapid determination of 16S ribosomal RNA sequences for phylogenetic analyses. Proc Natl Acad Sci. 1985; 82(20): 6966–6969.

22. Caporaso JG, Kuczynski J, Stombaugh J, Bittinger K, Bushman FD, Costello EK, et al. QIIME allows analysis of high-throughput community sequencing data. Nat Methods. 2010; 7:335–336. doi: 10.1038/nmeth.f.303 PMID: 2083131

23. Bragg L, Stone G, Imelfort M, Hugenholtz P, Tyson GW. Fast, accurate error-correction of amplicon pyrsequences using Acacia. Nat Methods. 2012; 9:425–426. doi: 10.1038/nmeth.1990 PMID: 22543370

24. Wu S, Zhu Z, Fu L, Niu B, Li W. WebMGA: a customizable web server for fast metagenomic sequence analysis. BMC Genomics. 2011; 12:445–456. doi: 10.1186/1471-2164-12-445 PMID: 21899761

25. Li W, Fu L, Niu B, Wu S, Wooley J. Ultrafast clustering algorithms for metagenomic sequence analysis. Brief in Bioinform. 2012; 13:656–668. doi: 10.1093/bib/bbs035

26. Caporaso JG, Bittinger K, Bushman FD, DeSantis TZ, Andersen GL, Knight R. PyNAST: a flexible tool for aligning sequences to a template alignment. Bioinform. 2010; 26(2): 266–267. doi: 10.1093/bib/btp636

27. Lozupone C, Lladser ME, Knights D, Stombaugh J, Knight R. UniFrac: an effective distance metric for microbial community comparison. ISME J. 2011; 5:169–172. doi: 10.1038/ismej.2010.133 PMID: 20827291

28. R core team. R: A language and environment for statistical computing. R foundation for statistical computing, Vienna, Austria. 2015; http://ww.R-project.org/.

29. Dixon P. VEGAN, a package of R functions for community ecology. J Veg Sci. 2003; 14(6): 927–930. doi: 10.1111/j.1654-1103.2003.tb02228.x

30. Champely S. PWR: Basic functions for power analysis. 2015; http://CRAN.R-project.org/package=pwr/

31. Levine TR, Hullett CR. Eta squared, partial eta squared and misreporting of effect size in communication research. Hum Commun Res. 2002; 28(4):612–625.

32. Schrader C, Schielke A, Ellerbroek L, Johne R. PCR inhibitors—occurrence, properties and removal. J Appl Microbiol. 2012; 113(5):1014–1026. doi: 10.1111/j.1365-2672.2012.05384.x PMID: 22747964

33. Niemi RM, Heikkila MP, Lahti K, Kalso S, Niemela SI. Comparison of methods for determining the numbers and species distribution of coliform bacteria in well water samples. J Appl Microbiol. 2001; 90 (6):850–858. PMID: 11412314

34. Tang JN, Shi XM, Shi CL, Chen HC. Characterization of a duplex PCR assay for the detection of entero- toxigenic strains of Staphylococcus aureus. J Rapid Methods Autom Microbiol. 2006; 14(3):201–217.

35. Rousselon N, Delgenes JP, Godon JJ. A new real time PCR (TaqMan PCR) system for detection of the 16S rRNA gene associated with fecal bacteria. J Microbiol Methods. 2004; 59(1):15–22. PMID: 15325749

36. Yu Z, Morrison M. Improved extraction of PCR-quality community DNA from digesta and fecal samples. Biotechniques. 2004; 36(5): 808–812. PMID: 15152600

37. Zhang BW, Li M, Ma LC, Wei FW. A widely applicable protocol for DNA isolation from fecal samples. Biochem Genet. 2006; 44(11–12): 503–512. PMID: 17094033

38. Willner D, Daly J, Whitey D, Grimwood K, Wainwright CE, Hugenholtz P. Comparison of DNA extraction methods for microbial community profiling with an application to pediatric bronchoalveolar lavage samples. PLoS One. 2012; 7:e34605. doi: 10.1371/journal.pone.0034605 PMID: 22514642

39. Deger G, Pakdemirili M, Candan F, Akgun S, Boyaci H. Strength of wheat and barley stems and design of new beam/columns. Math Comput Appl. 2010; 15(1):1–13.

40. Mackenzie BW, Waite DW, Taylor MW. Evaluating variation in human gut microbiota profiles due to DNA extraction method and inter-subject differences. Front Microbiol. 2015; 6:130. doi: 10.3389/fmicb.2015.00130 PMID: 25741335