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Strain-level metagenomic analysis of the fermented dairy beverage nunu highlights potential food safety risks

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Abstract

The rapid detection of pathogenic strains in food products is essential for the prevention of disease outbreaks. It has already been demonstrated that whole metagenome shotgun sequencing can be used to detect pathogens in food but, until recently, strain-level detection of pathogens has relied on whole metagenome assembly, which is a computationally demanding process. Here, we demonstrate that three short read alignment-based methods, MetaMLST, PanPhlAn, and StrainPhlAn, can accurately, and rapidly, identify pathogenic strains in spinach metagenomes which were intentionally spiked with Shiga toxin-producing Escherichia coli in a previous study. Subsequently, we employ the methods, in combination with other metagenomics approaches, to assess the safety of nunu, a traditional Ghanaian fermented milk product which is produced by the spontaneous fermentation of raw cow milk. We show that nunu samples are frequently contaminated with bacteria associated with the bovine gut, and worryingly, we detect putatively pathogenic E. coli and Klebsiella pneumoniae strains in a subset of nunu samples. Ultimately, our work establishes that short read alignment-based bioinformatics approaches are suitable food safety tools, and we describe a real-life example of their utilisation.

Importance

Foodborne pathogens are responsible for millions of illnesses, annually. Here, we demonstrate that short read alignment-based bioinformatics tools can accurately, and rapidly, detect pathogenic strains in food products from shotgun metagenomics data. The methods used here are considerably faster than both traditional culturing methods and alternative bioinformatics approaches that rely on metagenome assembly, and thus they can potentially be used for more high-throughput food safety testing. Overall, our results suggest that whole metagenome sequencing can be used as a practical food safety tool to prevent diseases or link outbreaks to specific food products.

Introduction

In recent years, high-throughput sequencing (HTS) has become an important tool in food microbiology (1). HTS enables in-depth characterisation of food-related microbial isolates,
via whole genome sequencing (WGS), and it facilitates culture-independent analysis of mixed microbial communities in foods, via metagenomic sequencing.

WGS has provided invaluable insights into the genetics of starter cultures (2, 3), and it is routinely used in epidemiology to identify outbreak-associated foodborne pathogens isolated from clinical samples, by comparing the single nucleotide polymorphism (SNP) profiles of outbreak strain genomes versus non-outbreak strain genomes (4-6). Metagenomic sequencing enables the elucidation of the roles of microorganisms during food production (7-9), and it can be used to track microorganisms of interest through the food production chain, as illustrated by Yang et al. (10), who used whole metagenome shotgun sequencing to track pathogenic species in the beef production chain. Indeed, metagenomic sequencing can be used to detect pathogens in foods to monitor outbreaks of foodborne illnesses (11), but few studies have done so, because of the limited taxonomic resolution achievable using these methods. Typically, 16S rRNA gene sequencing provides genus-level taxonomic resolution (12), and although sub-genus-level classification is achievable using species-classifiers (13) or oligotyping (14, 15), these methods cannot accurately discriminate between strains.

Similarly, metagenome sequence classification tools usually provide species-level resolution (16). However, strain-level resolution is necessary for the accurate identification of pathogens in food products (17). Leonard et al. successfully achieved strain-level resolution of Shiga toxin producing Escherichia coli strains in spinach samples using metagenome shotgun sequencing (18). However, the bioinformatics methods used in that study were based on metagenome assembly, which is a computationally demanding process (19, 20), and thus alternative strain-level identification methods are needed.

Since 2016, several short read alignment based software applications, including MetaMLST (20), StrainPhlAn (21), and PanPhlAn (19), have been released that can achieve strain-level characterisation of microorganisms from metagenome shotgun sequencing data. All three applications are considerably faster than metagenome assembly based methods. To date, these programs have not been employed to detect pathogens in food products, but there is strong evidence to suggest that they have considerable potential for this purpose: MetaMLST accurately predicted that the strain responsible for the 2011 German E. coli outbreak belonged to E. coli ST678 (20), and similarly, PanPhlAn accurately predicted that the strain was a Shiga toxin producer (19), based on the analysis of the gut metagenomes of infected patients (22). StrainPhlAn has so far not been used for epidemiological purposes, but a recent
study demonstrated that it can be used to predict the phylogenetic relatedness of bacterial strains from different samples (21).

MetaMLST aligns sequencing reads against a housekeeping gene database to identify sequence types present in metagenomic samples based on multilocus sequence typing (MLST). The MetaMLST database contains all currently known sequence types, but it can be updated as required to include newly identified sequence types. MetaMLST does not require any prior knowledge of the microbial composition of sample and it can simultaneously detect different species’ sequence types. PanPhlAn aligns sequencing reads against a species pangenome database, constructed from reference genomes, to functionally characterise strains present in metagenomic samples. PanPhlAn allows the user to generate customisable pangenome databases for any species. StrainPhlAn extracts species specific marker genes from sequencing reads and it aligns the markers against reference genomes to identify the strains present in metagenomic samples. StrainPhlAn requires output from MetaPhlAn2, and both programs use the same database.

In this study, we describe the characterisation of nunu, a traditional Ghanaian fermented milk product (FMP), at the genus, species, and strain-levels, using a combination of 16S rRNA gene sequencing and whole metagenome shotgun sequencing. Nunu is produced by the spontaneous fermentation of raw cow milk in calabashes or plastic or metal containers under ambient conditions, and it is usually consumed after 24-36 hours (23). At present, little is known about nunu’s microbiology, relative to other FMPs, like kefir or yoghurt (24). Previously, a number of potentially pathogenic bacteria, including 

Results

Enterobacter, Escherichia and Klebsiella, were detected in nunu by culture based methods (25). Here, we carry out the first culture-independent analysis of a number of nunu samples. In addition to detecting the presence of a variety of lactic acid bacteria (LAB) typical of fermented dairy products, MetaMLST, PanPhlAn and StrainPhlAn all indicated the presence of pathogenic E. coli and Klebsiella pneumoniae in a subset of the samples. We also demonstrate that these tools can accurately predict the presence of pathogenic strains in foods by testing them on food metagenomes which were spiked with Shiga toxin producing E. coli. Ultimately, our work establishes that short read alignment based methods can be used for the detection of pathogens in foods.
16S rRNA gene sequencing of nunu samples

Nunu samples were collected from producers with hygiene practice training (n=5) and producers without hygiene practice training (n=5), respectively. 16S rRNA gene sequencing analysis revealed that there were no significant differences in the alpha-diversity of nunu samples from trained or untrained producers (Figure S1a), although there was a clear separation in the beta-diversity of the two groups (Figure S1b).

The 16S rRNA data was also analysed to determine bacterial composition (Figure 1a). At the family level, all of the samples were dominated by Lactobacillales, and at the genus-level, most samples were dominated by Streptococcus, although the sample 1t2am was dominated by Lactococcus. Enterococcus was detected in 4/10 samples (1 trained and 3 untrained) at ≥3% relative abundance, and it was highest in the sample 2u6am, where it was present at 19% relative abundance. In addition, Staphylococcus was detected in all 10 samples, although its abundance was ≤1% in each case. The detection of staphylococci was consistent with a corresponding culture-dependent analysis of the samples (supplemental material).

Importantly, Enterobacteriales were also prevalent. Enterobacter was detected in 9/10 samples (4 samples from trained producers and 5 from untrained producers) at ≥1% relative abundance, and it was highest in the sample 2u8am, where it was present at 23% relative abundance. Escherichia-Shigella was detected in 8/10 samples (4 trained and 4 untrained) at ≥1% relative abundance, and it was highest in the sample 1t7am, where it was present at 17% relative abundance; this finding was again consistent with culture-dependent analysis of the samples (supplemental material).

The Kruskal-Wallis test indicated that there were significant differences in the relative abundances of Macrococcus (p=0.01), which was higher in samples from trained producers, and Streptococcus (p=0.02), which was higher in samples from untrained producers (Figure 1b). No other genera were significantly different.

Species-level compositional analysis of nunu samples as revealed by shotgun sequencing

MetaPhlAn2-based analysis of shotgun metagenomic data provided results that were generally consistent with those derived from amplicon sequencing. 11 species accounted for >90% of the microbial composition of every sample (Figure 2). At the species-level, most samples were dominated by Streptococcus infantarius, although sample 1t2am was
dominated by *Lactococcus lactis*. *Enterococcus faecium* was detected in 4/10 samples (2 trained and 2 untrained) at ≥1% relative abundance, and it was highest in the sample 1t2am, where it was present at 22% relative abundance. High abundances of *Enterobacteriaceae* were again apparent. *Enterobacter cloacae* was detected in the sample 1t8am, where it was present at 1% relative abundance. *Escherichia coli* was detected in 2/10 samples (2 trained) at ≥7% relative abundance, and it was highest in 1t7am, where it was present at 13% relative abundance. *Klebsiella pneumoniae* was detected in 7/10 samples (4 trained and 3 untrained) at ≥3% relative abundance, and it was highest in 1t8am, where it was present at 71% relative abundance. In contrast, *Klebsiella* was not detected by amplicon sequencing, and this discrepancy might be due to similarities in the 16S rRNA genes from these genera \(^{(42)}\).

The Kruskal-Wallis test indicated that there were significant differences in the relative abundances of *Macrococcus caseolyticus* (\(p=0.01\)), which was higher in samples from trained producers, and *Streptococcus infantarius* (\(p=0.01\)), which was higher in samples from untrained producers (Figure S2). No other species were significantly different.

**Investigation of the functional potential of the nunu microbiota**

SUPER-FOCUS was used to provide an overview of the functional potential of the nunu metagenome. As expected, a significant proportion of the metagenome was assigned to housekeeping functions like carbohydrate metabolism, nucleic acid metabolism, and protein metabolism (Figure 3). However, SUPER-FOCUS also detected high levels of functions associated with horizontal gene transfer and virulence in nunu. The level 1 subsystem “Phages, Prophages, Transposable elements” was present at ≥1% average relative abundance in both groups, although it was significantly higher in nunu samples from trained producers (\(p=0.047\)). Similarly, the level 1 subsystem “Virulence” was present at ≥3.5% average relative abundance in both groups.

HUMAnN2 was used to provide more comprehensive insights into the functional potential of the nunu metagenome. Unsurprisingly, the 25 most abundant genetic pathways were associated with carbohydrate metabolism, nucleic acid metabolism, and protein metabolism (Figure 4a). MDS analysis of all the normalised HUMAnN2 pathway abundances suggested that there were differences in the overall functional potential of the groups (Figure S3), and we detected significant differences in the relative abundances of some individual pathways (Table S1). Notably, we observed that histidine degradation pathways were higher in trained
samples (p=0.047) (Figure 4c). Furthermore, histidine decarboxylase genes were only detected in trained samples. Several other undesirable genetic pathways were detected in both groups. For example, putrescine biosynthesis pathways and polymyxin resistance genes co-occurred in 7/10 samples (Figure 4c), and these pathways were all attributed to *E. cloacae*, *E. coli*, *K. pneumoniae*, or a combination of these three species. We detected several other antibiotic resistance genes, including beta-lactamase genes and methicillin resistance genes, in both groups (Figure S4). In addition, we found HGT-associated genes, including plasmid maintenance genes and transposition genes, in both groups.

### Application of strain-level analysis to characterise enteric bacteria in nunu

Leonard *et al.* previously used metagenomic sequencing to detect *E. coli* in spinach which was intentionally spiked with *E. coli* O157:H7 strain Sakai (11). We downloaded the metagenomic reads from that study (16 samples) and we subjected them to StrainPhlAn, MetaMLST and PanPhlAn analysis, to confirm that these tools can accurately detect pathogens in food samples: MetaMLST was used for multi-locus sequence typing, StrainPhlAn was used for phylogenetic identification, and PanPhlAn was used for functional characterisation. MetaMLST accurately detected *E. coli* ST11 in 7/16 spinach samples (Table 1). StrainPhlAn detected *E. coli* strains in 5/16 samples and it showed that the *E. coli* strain in each of these samples was closely related to *E. coli* O157:H7 strain Sakai (Figure 5). PanPhlan detected Shiga toxin genes in 15/16 samples (Table 1) and it indicated that the *E. coli* strain in each of these samples was most closely related to *E. coli* O157:H7 strain Sakai. Thus, overall, PanPhlAn was the most sensitive method in this instance, since it was able to detect STEC in almost all of the samples, whereas the other tools detected STEC in less than half of the samples. In a follow-on study, Leonard *et al.* spiked spinach with 12 different Shiga toxin producing *E. coli* strains, and they detected single strains in 17 samples (18). We downloaded the metagenomic reads from the 17 samples and ran PanPhlAn, and were able to identify Shiga toxin genes in all 17 samples (Table S2).

Having established the relative merits of these tools, we subsequently employed all three strategies to identify the strains of *E. coli* and *K. pneumoniae* present in the nunu samples. With regard to *E. coli*, MetaMLST detected a novel *E. coli* sequence type in 17 strains (Table 2). StrainPhlAn detected 24 *E. coli* marker genes in the samples and a phylogenetic tree (Figure 6a), which was generated by aligning these markers against 118 *E. coli* reference genomes.
listed in Table S3), revealed that the *E. coli* strain in one sample, 1t7am, was closely related to *E. coli* O139:H28 E24377A. PanPhAn detected *E. coli* strains in two samples: 1t7am and 1t8am. MDS analysis indicated that the strains from the two samples were functionally distinct from one another. Notably, a ShET2 enterotoxin encoding gene was identified in the *E. coli* strain from 1t7am. The same gene was found in *E. coli* O139:H28 E24377A. With regard to *K. pneumoniae*, MetaMLST detected the known sequence type *K. pneumoniae* ST39 in the sample 2u3am. Apparently novel *K. pneumoniae* sequence types were identified in six other samples (Table 1). StrainPhlAn detected 38 *K. pneumoniae* marker genes in the samples and a phylogenetic tree (Figure 6b), which was constructed by aligning these markers against 40 *K. pneumoniae* reference genomes (listed in Table S4), revealed that the *K. pneumoniae* strains in two samples, 1t8am and 2u3am, were closely related to *K. pneumoniae* KpQ3. In contrast, the *K. pneumoniae* strain in 1t7am was most closely related to *K. pneumoniae* UCICRE 7. MDS analysis of the PanPhlAn output showed that five of the detected *K. pneumoniae* strains were functionally similar to one another (Figure 6c). However, two of the detected *K. pneumoniae* strains, in samples 1t6am and 1t7am, appeared to be functionally distinct from the others. In addition, PanPhlan indicated that sample 1t6am might have contained multiple strains, since an unusually high number of 5746 *K. pneumonia* gene families were detected. A TEM beta-lactamase gene was found in 1t2am using PanPhlAn and, furthermore, an OXA-48 carbapenemase gene was detected in 2u8am and the same gene was found in *K. pneumoniae* KpQ3.

Finally, we compared the time taken to process 10 nunu metagenome samples using the short-read alignment tools versus the metagenome assembler IDBA-UD (Figure S5). In each case, we observed that all of the short-read alignment tools were faster than IDBA-UD. It is important to note that additional bioinformatics analyses (contig binning, SNP analysis, etc.) are required to achieve strain-level identification from assembled metagenomes, and this emphasises the superior speed of the short-read alignment tools.

**Discussion**

Foodborne pathogens are responsible for millions of cases of disease annually, in the United States alone (43). High-throughput sequencing can potentially be used to detect pathogenic strains in food products to prevent the occurrence of disease outbreaks. A recent proof of concept study demonstrated that whole metagenome shotgun sequencing accurately detected
Shiga toxin producing \textit{E. coli} (STEC) strains in spiked spinach samples (18). However, that study used whole metagenome assembly-based approaches to achieve strain-level taxonomic resolution of the STEC in the samples. Whole metagenome assembly is a computationally intensive, time-consuming process, as illustrated by Nurk \textit{et al.}, who recently reported that metagenome assembly can take between 1.5 hours to 6 hours, with a memory footprint ranging from 7.3 GB to 234.5 GB, to process a single human gut metagenomic sample, depending on the chosen assembler (44). Thus, the application of more rapid, less intensive bioinformatic tools for strain detection is desirable. In this study, we demonstrate that the short read alignment-based programs MetaMLST, StrainPhlAn, and PanPhlAn can accurately identify pathogens in food products.

We validated the accuracy of each approach by processing spinach metagenome data from samples that were spiked with the STEC O157:H7 Sakai in a previous study (11). We observed that PanPhlAn was the most sensitive approach. Indeed, PanPhlAn was able to identify STEC in every sample where it was present at >2% relative abundance, whereas the other approaches worked best when STEC was present at high relative abundances. However, none of the tools detected \textit{E. coli} O157:H7 Sakai in every sample tested. The observation of false negatives highlights that the tools are not entirely accurate. It is likely that increased sequencing depth and/or longer sequencing read lengths would reduce the false negative rate.

We recommend that these tools be used to supplement data from metagenome sequence classifiers like MetaPhlAn2, which did detect \textit{E. coli} in each sample. Therefore, we subsequently used the strain-level analysis tools in combination with other metagenomic approaches to assess the safety of nunu, a traditional Ghanaian fermented milk product.

Nunu is produced through the spontaneous fermentation of raw cow milk in calabashes or other containers for 24-36 hours at ambient temperature (23). The crude nature of the nunu production process has raised food safety concerns (25). Indeed, several potentially pathogenic microorganisms were previously detected in nunu samples by microbial culturing (25). This resulted in some nunu producers receiving hygiene practice training to improve food safety. However, our work suggests that there is little difference in the prevalence of pathogens in nunu samples from trained and untrained producers. One reason for this may be that it is difficult for the nunu producers to adhere to the training recommendations which are not appropriate to the rural production conditions. During training, the producers were advised to pasteurise the milk before cooling and adding a starter culture. After incubating for 4-6 hours in a covered container, they were advised to stir the mixture and refrigerate the...
product. Lack of access to specific heat control and electricity, as well as the variance from the traditional method, which does not use a starter culture, are both reasons why the training is not adhered to.

16S rRNA gene sequencing revealed that the samples were dominated by Lactobacillales. However, we also detected high abundances of Enterobacteriales, including Enterobacter and Escherichia, in both groups. Subsequently, whole metagenome shotgun sequencing showed that most samples were dominated by Streptococcus infantarius, a species which was previously identified in other African dairy products (45, 46). Concernedly, S. infantarius has been linked to several human diseases, including bacteraemia (47), endocarditis (48) and colon cancer (49). Aside from S. infantarius, two other potentially pathogenic species, Escherichia coli and Klebsiella pneumoniae, were identified in a subset of samples.

Overall, our findings indicate that nunu samples from trained producers and untrained producers were contaminated with faecal material. Cattle faeces can be a major source of bacterial contaminants in raw cow milk (29), and thus, our results are not entirely surprising, but the remarkable abundance of such microorganisms in nunu is worrying. It had been hoped that nunu could be used to supplement traditional cereal-based weaning foods to improve infant nutrition. However, qualitative research among mothers and health workers highlighted safety concerns, which, as we have shown here, are valid. In particular, the presence of E. coli and K. pneumoniae in nunu is a concern, and, thus, we employed strain-level metagenomics for the further characterisation of these bacteria.

In terms of E. coli, strain-level analysis indicated that the E. coli strain in one sample was an enterotoxin producer and it was closely related to E. coli O139:H28 E24377A, a strain which was linked to an outbreak of waterborne diarrhoea in India (50). In terms of K. pneumoniae, strain-level analysis indicated that the K. pneumoniae strains in two samples were antibiotic resistant and they were closely related to K. pneumoniae KpQ3, a strain which was linked to nosocomial outbreaks among burn unit patients. Thus, strain-level analysis suggests that there are likely pathogens in some of the samples. Interestingly, PanPhlAn also suggested that there were functionally distinct strains of both species in nunu samples from different producers. Perhaps, this indicates multiple incidences or sources of contamination.

Undoubtedly, our work highlights an urgent need to further improve hygiene practices during nunu production, and the pasteurisation of the starting milk and the use of starter-based fermentation systems is an obvious solution.
In conclusion, our work suggests that short read alignment-based strain detection tools can be used to detect pathogens in other foods, apart from nunu or spinach, and they might also be useful for tracing the sources of foodborne disease outbreaks back to particular foods. Such tools are a significant improvement over 16S rRNA gene sequencing, which is often limited to genus-level identification, or metagenome read classification tools, which are limited to species-level identification (16). In addition, they are faster, and less computationally intensive, than metagenome assembly-based strain detection methods, making them more relevant to real-life scenarios which necessitate the rapid testing of many food samples. With DNA sequencing costs continuing to decrease, the approach outlined here is an affordable option for food safety testing.

Acknowledgements

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Materials and Methods

Sampling

Five nunu samples were collected from producers with hygiene practice training, and another five samples were collected from producers without hygiene practice training. The identity of the samples from trained and untrained individuals was blinded until after sequencing analysis was completed. The samples from the trained group were labelled 1t2am, 1t6am, 1t7am, 1t8am, and 2t2am. The samples from the untrained group labelled 1u6am, 2u2am, 2u3am, 2u6am, and 2u8am. All samples were collected in the morning and placed on ice for transport to the lab. Sample aliquots (4ml) were then mixed with glycerol to a final concentration of 20% and stored at -20°C prior to DNA extraction. DNA was extracted from
the samples at the Animal Research Institute, Accra, Ghana and then sent to Scotland to comply with International laws on the import of animal samples (Import Licence form AB117).

Microbiological analysis

Basic microbiology culture analysis was carried out in Ghana. The plate-count technique was used to estimate the total viable bacterial count of the nunu samples on Milk Plate Count Agar (LAB M, UK). Bacterial counts were compared for plates growing aerobically or anaerobically at 30°C for 36-72 h. Anaerobic plates were incubated in airtight canisters containing CO₂:Gen sachets (Oxoid, UK), which created an anaerobic atmosphere. Following incubation, colonies were counted using an SC6+ electronic colony counter (Stuart Scientific, UK). The presence of specific pathogens in the nunu samples was determined by streaking nunu directly onto selective agar plates to visually assess bacterial growth. The following selective agars were used: Blood agar (Merck, Germany) for Staphylococcus; MacConkey agar (Merck, Germany) for Enterobacteria; de Man Rogosa Sharpe agar (MRS) (Oxoid, UK) for Lactobacillus species; and Salmonella Shigella agar (Oxoid, UK). Any mixed growth plates were re-purified by streaking onto selected secondary agars. Lactose fermenting colonies identified on MacConkey agar were sub-cultured onto Eosin Methylene Blue Agar (EMBA) (Scharlau Chemie, Spain) to isolate/identify E. coli. Additionally, Staphylococcus colonies from Blood Agar were sub-cultured onto Mannitol Salt Agar (MSA) (Oxoid, UK) to isolate/identify Staphylococcus aureus. The following biochemical tests were used to confirm bacterial identification: the Motility Indole Urea (MIU) test; the catalase test; the Triple Sugar Iron (TSI) test; and the Indole Methyl Red Vorges-Prosekur Citrate (IMViC) tests.

Cellular morphology was determined by Gram staining as well as microscopic examination.

DNA extraction and next generation sequencing

Briefly, 1 ml of each thawed sample was diluted in 9 ml of sterile PBS, mixed thoroughly using vortex and centrifuged for 10 min (8,000-10,000 g). The bacterial cell pellets were resuspended in 432 µl sterile dH₂O and 48 µl 0.5 M EDTA, mixed thoroughly by a combination of vortex and with a sterile pipette tip and the suspension frozen. The frozen samples were thawed on the bench and refrozen and finally thawed (giving a total of two
freeze/thaw cycles) before extracting the DNA using the Promega Wizard genomic DNA extraction kit (Promega, Madison, WI, USA) according to the manufacturer’s protocol. The freeze/thaw cycles were carried out to maximise bacterial cell lysis. Following extraction, the DNA pellets were air dried for about 60 minutes and stored sealed under airtight conditions and transported from the Animal Research Institute, Accra, Ghana to the Rowett Institute, at University of Aberdeen, for further analysis.

DNA extracts were quantified using the Qubit High Sensitivity DNA assay (BioSciences, Dublin, Ireland). 16S rRNA gene sequencing libraries were prepared from extracted DNA using the 16S Metagenomic Sequencing Library Preparation protocol from Illumina, with minor modifications (26). Samples were sequenced on the Illumina MiSeq in the Teagasc sequencing facility, with a 2 x 250 cycle V2 kit, in accordance with standard Illumina sequencing protocols. Whole-metagenome shotgun libraries were prepared in accordance with the Nextera XT DNA Library Preparation Guide from Illumina (26). Samples were sequenced on the Illumina MiSeq in the Teagasc sequencing facility, with a 2 x 300 cycle V3 kit, in accordance with standard Illumina sequencing protocols.

**Bioinformatics**

Raw 16S rRNA gene sequencing reads were quality filtered using PRINSEQ (27). Denoising, OTU clustering, and chimera removal were done using USEarch (v7-64bit) (28), as described by Doyle et al. (29). OTUs were aligned using PyNAST (30). Alpha-diversity and beta-diversity were calculated using Qiime (1.8.0) (31). Taxonomy was assigned using a BLAST search (32) against SILVA SSU 119 database (33).

Raw whole-metagenome shotgun sequencing reads were filtered, on the basis of quality and quantity, and trimmed to 200 bp, with a combination of Picard Tools (https://github.com/broadinstitute/picard) and SAMtools (34). MetaPhlAn2 was used to characterise the microbial composition of samples at the species-level (35). MetaMLST (20), PanPhlAn (19), and StrainPhlAn (21) were used to characterise the microbial composition of the samples at the strain-level. GraPhlAn (36) was used to construct phylogenetic trees from the StrainPhlAn output. SUPER-FOCUS (37) and HUMAnN2 (38) were used to determine the microbial metabolic potential of samples. IDBA-UD (39) was used for metagenome assembly.
Accession numbers

Sequence data have been deposited in the European Nucleotide Archive (ENA) under the project accession number PRJEB20873.

Statistical analysis

Statistical analysis was done in R-3.2.2 (40). The Kruskal-Wallis test was done using the compareGroups package, and the resulting p-values were for multiple comparisons. PCoA analysis of 16S rRNA gene sequencing data was done using the phyloseq package (41). Multidimensional scaling (MDS) was done using the vegan package. Data visualisation was done using the ggplot2 package.

References


enterotoxigenic Escherichia coli strain E24377A, obtained from a tribal drinking water source in India. Genome announcements 3:e00225-00215.
Table 1. The results of MetaMLST and PanPhlAn analysis of spinach metagenomes spiked with *E. coli* O157:H7 Sakai

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Table 2. The results of MetaMLST analysis of the nunu metagenomic samples

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Figure legends

Figure 1. 16S rRNA gene sequencing based analysis of nunu samples. (A) Heat map showing the 25 most abundant bacterial genera across the nunu samples. (B) Bar plot showing genera which were differentially abundant in either group.

Figure 2. The species-level microbial composition of nunu samples, as determined by MetaPhlAn2.

Figure 3. The average abundances of the SUPER-FOCUS Level 1 functions that were detected in nunu samples.

Figure 4. HUMAnN2 analysis. (A) Heat map showing the 25 most abundant MetaCyc pathways detected across the ten nunu metagenomic samples. (B) Bar plot showing differences in histidine metabolic potential between nunu samples from trained producers and nunu samples from untrained producers. (C) Bar plots showing the relative contributions of E. cloacae, E. coli and K. pneumoniae to the MetaCyc pathways PWY-6305 (putrescine biosynthesis) and PWY0-1338 (polymyxin resistance).

Figure 5. StrainPhlAn analysis of the spinach metagenome.

Figure 6. Strain-level analysis. Phylogenetic trees showing the relationships between (A) E. coli strains and (B) K. pneumoniae strains detected in the nunu metagenomic samples and their respective reference genomes, as predicted by StrainPhlAn. (C) MDS showing the functional similarities between strains detected in the nunu metagenomic samples, as predicted by PanPhlAn; reference genomes are shown in faded grey.
StrainPhlAn: E. coli (spinach metagenome)