In vitro growth of gut microbiota with selenium nanoparticles

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A B S T R A C T

The application of nanoparticles rose steeply in the last decade, where they have become a common ingredient used in processed human food, improving food properties such as shelf life and appearance. Nanoparticles have also attracted considerable interest to the livestock industry, due to their efficacy in intestinal pathogen control, with the regulatory and consumer driven push for the removal of antibiotic growth promoters. The influence of selenium (Se) nanoparticles was investigated on a diverse and mature broiler caecal microbiota using in vitro culturing and 16S rRNA gene sequencing methods for microbiota characterisation. Caecal microbiota was collected from 4 traditionally grown heritage roosters and grown for 48 h, in the presence and absence of Se nanoparticles, with 2 technical replicates each. The effect of rooster as a biological variable strongly overpowered the effects of nano-Se in the media, resulting in moderate effects on the structure and diversity of the caecal microbial community. However the nanoparticles showed a significant reduction (P < 0.05) in the abundance of an emerging poultry pathogen, Enterococcus cecorum identical operational taxonomic units (OTU), which could be of notable interest in poultry production for targeted E. cecorum control without significant disturbance to the total microbial community.

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1. Introduction

The application of nano-scaled materials, 1 to 100 nm, has rapidly expanded across various disciplines, including but not limited to, electronics, technology, consumer goods, biomedical science, agriculture and microbiology. They display unique physicochemical properties due to high surface energy and increased surface area to volume ratio (Regan et al., 2012). Nanoparticles are used in everyday applications such as self-cleaning surfaces (Mueller and Nowack, 2008), topical products (Goyal et al., 2016) and food preservation (Espitia et al., 2012). They have excellent antimicrobial properties through the disruption of microbial cell membranes (Hajipour et al., 2012; Thill et al., 2006) and oxidation (Le Ouay and Stellacci, 2015). Culture studies have been used to examine their biocidal interaction (Jia et al., 2017; Teodor et al., 2011) towards both prokaryotic, bacterial pathogens and eukaryotic cells such as tumour and stem cells (Arora et al., 2008; Greulich et al., 2009; Kaul and Amiji, 2005). They have also been intensively investigated for their use in joint and bone reconstruction therapies (Gangadoo et al., 2015), agricultural products (Gangadoo et al., 2016) and delivery of drugs and other substances to the body (Gupta and Curtis, 2004), as they exhibit high biocompatibility (Lu et al., 2010; Naahidi et al., 2013) and biodegradability (Mahapatro and Singh, 2011; Panyam and Labhasetwar, 2003).

Nanoparticles can be used as vehicles to transport substances to the body effectively and fast, by avoiding complex pathways and defence mechanisms as compared to their bulk counterparts (Desai et al., 1996; Mohanraj and Chen, 2006). Many studies have shown
the positive effect of NP formulations delivered to the gut microbiota, focussing primarily on reducing the pathogenic load with an antibiotic based approach, by inhibiting the growth of harmful microbes (Karavolos and Holban, 2016). Since various materials have successfully targeted detrimental microbes through NP delivery, it was proposed that a NP-based system, using metal salts and complex, could also be used to enhance beneficial bacteria by delivering essential nutrients to the gut microbiota. Nanoparticles have been used to increase the abundance of beneficial species such as *Lactobacillus* and *Bifidobacteria* and reduce pathogenic bacteria and coliform counts (Gangadoo et al., 2018; Han et al., 2010; Yausheva et al., 2018). The gut ecology is a complex community and it is necessary to consider the complex interactions of multiple bacterial species, the chemistry of their growth environment and the metabolites produced.

Selenium (Se) is an important trace element required by the body for the proper functioning and development of the immune system, and it is routinely supplemented to poultry rations to prevent detrimental effects of Se deficiency in birds (Gangadoo et al., 2016). About one quarter of the gut microbiome has the ability to express selenoproteins and Se availability in microbiological media affects their expression (Kasaikina et al., 2011). These proteins play an important role in both bacteria and mammalian host where they are essential in numerous bodily functions (Labunsky et al., 2014). We have previously investigated the ability of selenium nanoparticles (nanoSe) to improve the delivery of Se to birds and have characterised the resulting modifications of the intestinal microbiota (Gangadoo et al., 2018). We found increased abundance of some beneficial bacteria, for example *Lactobacillus* sp. and *Faecalibacterium prausnitzii*, however, without significant pathogen reduction. The quantity of butyric acid in different gut sections was increased. Butyric acid is a primary energy source for intestinal colonocytes and can promote good gut health (Van Immerseel et al., 2017).

Traditionally raised free range chickens generally show higher diversity of intestinal microbiota compared to intensively reared birds (Chen et al., 2018; Cui et al., 2017). Their microbiota may be used to increase the abundance of beneficial bacteria, for example *Lactobacillus plantarum* (ATCC BAA-793) and *Lactobacillus rhamnosus* (ATCC 53103) to mid-stationary phase in LYHBHI media. The supernatant of the bacterial ferments were then mixed at a 1:1 ratio with a final volume of 50 mL, filter sterilised and added to the media. The volume of water was adjusted to ensure that the original LYHBHI medium was not diluted. The enriched LYHBHI was purged for 30 min prior to inoculation of caecal content with anaerobic gas mix (80%N2/10%CO2/10%H2, BOC, Queensland, Australia).

NanoSe was prepared as previously described (Gangadoo et al., 2017). Briefly, selenium tetrachloride was reduced with ascorbic acid to Se atoms, to which a protecting agent, polystyrene-4-sulfonate, was added to allow the formation of nanoparticle clusters. The synthesised, dark red solution was washed by multiple centrifugation with Milli-Q water and a full characterisation, including size, shape, morphology and crystallinity, was conducted. The nanoSe was then diluted with Milli-Q water to 0.9 mg/kg.

2.3. Cecum starter cultures

Caeca, from 4 roosters, were donated by a local heritage breeder. The caeca were raised with organic feed without antibiotics and had exclusive outdoor access, including overnight outdoor roosting, which provided intensive contact with wild flora and fauna. The whole intestine of each rooster was removed and placed immediately into an anaerobic gas pack (Cat. #260683, BD GasPak EZ Pouch Systems) and stored at –20 °C. The caecal samples were slowly allowed to defrost at 4 °C for 30 min. The contents of the whole caeca, for each rooster separately, were squeezed out and diluted in 50 mL of enriched LYHBHI media with 15% glycerol in an anaerobic work station (A35, Whitley, Shipley, UK). The caecal starter cultures for each rooster’s caecal content were then aliquoted as 50 x 1 mL stock and stored at –80 °C until the start of the experiment. This would eliminate cold sensitive species and allow the reproducible use of each 1 mL stock for the future in vitro experiments.

2.4. In vitro growth cultures

On the day of the experiment, a single glycerol stock for each one of the 4 roosters was thawed and inoculated into 50 mL of enriched LYHBHI media to grow parent cultures for the experimental inoculation. The experimental cultures were prepared in 20 mL of media in 50 mL Erlemeyer flask with a cotton stopper, allowing for gas exchange, and incubated at 37 °C on a digital orbital shaker (Heathrow Scientific), shaking at a speed of 0.21 × g in an anaerobic hood (Whitley A35 Anaerobic Workstation, UK).
running on a nitrogen rich gas mix (80% N2/10% CO2/10% H2). Four cultures were prepared from each rooster’s caecal content, 2 as control and 2 with 0.9 mg/kg of nanoSe, by inoculating late exponential growing parental culture to achieve a starting culture OD620 of 0.1. Thus, the final experiment was performed on 16 cultures; n = 8 for control and the nanoSe treatment each, on 4 biological replicates (rooster’s caecal content) and 2 technical replicates each, as shown in Fig. 1. Sampling of the cultures was done at 24 and 48 h and the samples were centrifuged at 18,500 × g at 4 °C for 10 min. The pellets and the supernatants were used for microbial and metabolite analysis, respectively.

2.5. DNA extraction

DNA was extracted from the centrifuged pellets of the microbial cultures. The lysis step was based on the method suggested by Yu and Morrison (2004), followed by a DNA purification step. The lysis buffer (0.5 mL) and 0.1 g of sterile zirconia beads (Cat. #11079101, BioSpec Products) were added prior to bead-beating (Mini-beadbeater, BioSpec Products) for 5 min. Following a 15-min incubation at 85 °C, the samples were centrifuged for 5 min, and binding buffer (0.8 mL) was added to the supernatant and placed through DNA Silica Membrane Mini Spin Column (Cat. #1920-250, Epoch Life Science, Inc.), followed by a two-step washing with wash buffer (0.7 mL). The washed and dried column was then eluted with 50 μL of elution buffer. The composition of the buffers is included in Appendix Table 2.

2.6. DNA amplification and sequencing

Sequencing of 16S rRNA gene DNA amplicons was performed on the Illumina MiSeq platform using 2 × 300 bp paired-end sequencing. Primers were selected to amplify the V3 – V4 region of 16S rRNA genes: forward 5′-ACTCCTACGGGAGGCAGCAG-3′ and reverse 5′-GGACTACHVGGGTWTCTAAT-3′. The primers contained barcodes, spaces and Illumina sequencing linkers as previously described (Fadrosh et al., 2014). Two samples, one from the rooster 2 and one from the rooster 4, failed the sequencing process, and were thus excluded from the analysis.

2.7. Statistical analysis

The analysis of microbial communities was performed in QIIME v1.9.1 (Caporaso et al., 2010). Paired end sequences were joined using the Fastq-Join algorithm and no allowed mismatches using only sequences with Phred quality threshold higher than 20. Operational taxonomic units were picked at 97% similarity using Uclust (Edgar, 2010) and inspected for chimeric sequences using Pintail (Ashelford et al., 2005). Taxonomic assignments were performed against the GreenGenes (DeSantis et al., 2006) database and QIIME default arguments. Further data exploration was done using Calypso (Zakrzewski et al., 2016). Total sum normalisation and a square root transformation was performed prior to statistical analysis. Student’s t-test was used to detect the significance of the differences between the groups. Analysis of similarities (ANOSIM) was performed using Calypso on weighed and unweighed UniFrac distance matrices calculated in QIIME, each with 99,999 permutations. Calypso was also used to implement the supervised multivariate Redundancy Analysis (RDA) using 999 permutations and linear regression analysis using Pearson correlation.

The complete annotated sequence dataset is publicly available on the MG-RAST database under library ID (Id pending).

2.8. Short-chain fatty acid analysis

The supernatants from the caecal cultures were diluted with 70% ethanol, filtered through a 0.45 μm syringe filter (Cat. #54504-RC, ThermoFisher) and analysed on the Gas Chromatography-Mass Spectrometry (GCMS) system. A standard stock solution (100 mg/kg) was used to construct calibration curves and stored as a method processing parameter in scan mode for the following short-chain fatty acids (SCFA), acetic, n-butyric, isobutyric, propionic and n-valeric acid.

The GCMS system used for metabolite analysis was a Shimadzu QP2010-Plus, fitted with a high-polarity column SH-Rxi-5Sil-MS (30 m × 0.25 mm × 0.25 μm, Restek) and equipped with an AOC-201 i.s autosampler. The GC oven temperature programme started at 100 °C and was held for 1 min, increased to 12 °C per min to a temperature of 170 °C, and ramped at 100 °C per min, until a final temperature of 260 °C was reached and held for 1 min (a total of an 8.73-min programme). The GC oven temperature was set as presented in Appendix Table 3. A sample of 1 μL was injected at 250 °C using helium (5.0, Coregas, Australia) as a carrier gas at 1.97 mL/min in a split injection mode. The pressure was held at 143.3 kPa, with a total He flow of 103.4 mL/min and using a split ratio of 5. The mass spectrometer was operated in the electron ionisation mode at 0.2 kV with a source temperature of 220 °C where scan mode was used from 33 to 150 m/z. The peaks were identified by matching the mass spectra with the National Institute of Standards and Technology (NIST) library, http://chemdata.nist.gov/.

![Fig. 1. A schematic of the in vitro experiment performed to examine the effect of nanoSe on growth cultures of rooster caeca samples.](image-url)
3. Results

3.1. Sample origin influences overall microbiota composition and abundance

The origin of caeca greatly influenced the microbiota community of the samples, showing great biological variation. The abundance of phyla *Firmicutes*, *Bacteroidetes*, *Proteobacteria* and *Actinobacteria* differed significantly between the 4 roosters (ANOVA, $P < 0.001$). *Lactobacillus* (>20%) was the dominant genus in all roosters combined, followed by *Streptococcus* (>15%), *Enterococcus* and *Clostridium* (>5%) (Appendix Table 4). Principal coordinates analysis (PCoA), as shown in Fig. 2A, was performed on both unweighed and weighed UniFrac matrices and shows similarities between roosters 2 and 3, while roosters one and 4 were very distinctive (Fig. 2B). The microbiota of roosters 2 and 3 included multiple genera not found in roosters one or 4, such as *Collinsella*, *Coprobacillus*, *Slackia*, and unclassified families of *Burkholderiales* and *Ruminococcaceae*. Rooster 4 had the most distinctive microbiota; dominated by *Clostridium*, with lower abundance of *Lactobacillus* compared to the other roosters, and higher amounts of *Trichococcus*, *Proteus* and unclassified families comprising of *Clostridiales* and *Burkholderiales*.

3.2. NanoSe influence on microbial composition and metabolite production

The enriched LYHBHI supported the growth of a diverse range of genera comprising of multiple previously uncultured species as shown in Appendix Table 3. NanoSe supplementation significantly ($P < 0.05$) increased 20 operational taxonomic units (OTU), as shown in Fig. 3A and reduced 8 OTU ($P < 0.05$), one of which was identified as 100% identical across the amplified region to *Enterococcus cecorum*, followed by 2 other *Enterococcus* OTU significantly reduced by nanoSe (Fig. 3B). *Enterococcus* OTU, including pathogenic *E. cecorum*, were exclusively reduced by nanoSe while genera *Lactobacillus* and *Streptococcus* were rearranged with some OTU significantly reduced and other significantly increased. Although nanoSe treatment was correlated with changes in abundance of some specific OTU an ANOSIM multivariate analysis of group similarities showed that the overall gut microbial composition was not affected by nanoSe ($P = 0.991$) or an additional 24 h of incubation.

![Fig. 2](image-url) Gut microbiota profile was clustered based on the bird sample. (A) PCoA analysis performed on weighed UniFrac matrices shows gut community profiles clustered by bird origin. (B) The multiple genera present in the samples confirm similarities and differences between rooster’s gut communities. PCoA – Principal coordinates analysis.
Furthermore, alpha diversity indices (Shannon’s index, richness and evenness) were also not affected by nanoSe or time of incubation ($P > 0.05$). The supplementation of nanoSe had no effect (using $t$-test) on SCFA production, as shown in Fig. 3C.

### 3.3. Interaction between gut community and short-chain fatty acid

PERMANOVA showed that SCFA production was significantly related to the microbiota composition ($P = 0.00067$) and RDA demonstrated that the microbial composition was significantly related to the SCFA ($P < 0.01$). The 5 SCFA, acetic acid, butyric acid, isobutyric acid (IBA), propanoic acid and valeric acid correlated with the abundance of a number of taxa (Fig. 4A). Valeric acid and IBA strongly correlated ($P < 0.001; R > 0.85$) with the same genera, including *Adlercreutzia*, *Desulfovibrio*, *Microbacterium*, unclassified Barnesiellaceae, unclassified Helicobacteraceae and unclassified WPS2, (Fig. 4B). Butyric acid and acetic acid shared one genus, an unclassified Clostridiales, exhibiting a strong correlation ($P < 0.001; R > 0.85$). Butyric acid additionally had a positive correlation ($P < 0.001; R = 0.86$) with *Clostridium* and an inverse correlation ($P < 0.001; R = -0.90$) with an unclassified Streptococcaceae.
4. Discussion

The human and animal microbiota is continuously altered with different lifestyles and environmental changes, and has undergone major rearrangements since the introduction of industrialised, large-scale food production in the last few centuries (Flandroy et al., 2018). This change in eating habits and the subsequent changes in gut microbiota has led to the modern age being described as an age of “microbiota genocide” (Sonnenburg et al., 2016). The lifestyle and eating habits of hunter-gatherer societies are very different to that of modern western societies and the difference drive profound changes when comparing ancient and modern human microbiotas (Davenport et al., 2017; Kumar and Forster, 2017; Warinner et al., 2015). This effect spills over to livestock and birds with characteristics microbiota changes occurring because of altered husbandry and feeding practices. Industrial scale grown birds experience very different growth conditions compared to their ancestors; the eggs are hatched under highly clean conditions, removing the influence of parental microbiota passage to the next generation (Donaldson et al., 2017). This results in aberrant microbiotas and high microbiota variation from one batch of hatchlings to another (Stanley et al., 2013). Microbiota analyses of chicken caeca across various...
projects have displayed an enormous discrepancy between bacterial species present in industrial birds and those present in birds grown in traditional low density open housing ways such as that found with “village chickens” or “backyard chickens” as called in Australia. Here we used the caeca of backyard chickens to investigate the effects of Se nanoparticles (nanoSe) on gut microbiota. The gut microbiota of an industrially grown domestic chicken, Gallus gallus domesticus, is typically comprised of 4 main phyla; Firmicutes, Bacteroidetes, Proteobacteria and a low amount of Actinobacteria (Oakley et al., 2014; Waite and Taylor, 2014; Wei et al., 2013). The high number of unclassified genera, presented in this study, is possibly indicative of the influence of non-industrialised housing and other environmental conditions (Kers et al., 2018), such as access to pasture, live plant and insect food content, full free range, and exposure to wild birds and animals.

Culturable genera, Lactobacillus, Streptococcus, Clostridium and Enterococcus strongly dominated (>80%) the rooster’s caecal community, while numerous uncultured genera remained in low abundance. The 80% to 90% sequence similarities render it impossible to infer function, pathogenicity or probiotic potential of these unidentifiable species to known cultured bacteria. The unknown and uncultured species often require metabolic feedback from other bacteria and can be cultured only in a complex community rather than as a single culture. The caecal microbiota communities were more diverse and different to the ones previously investigated with live birds treated with different concentrations of nanoSe (Gangadoo et al., 2018), and consequently the in vitro response of cultured caecal microbiota to nanoSe proved dissimilar to that seen in the microbiota of treated birds, including the lack of SCFA and Lactobacillus genus stimulation. It is not clear whether the different test systems or the different starting microorganisms have more pronounced influence in producing the different outcomes.

In contrast, the reduction of an emerging avian pathogen, E. cecorum, and 2 unknown enterococcus species, was observed with nanoSe at a concentration as low as 1 mg/kg. E. cecorum has been linked to enterococcal spondylitis and femoral head necrosis, resulting in symptoms such as hind limb weakness (Borst et al., 2017; Dolka et al., 2016) and lameness in poultry (McNamee and Smyth, 2000). Other symptoms observed include arthritis and spinal lesions (Dolka et al., 2017), with E. cecorum infection leading to a marked increase in flock mortality among all poultry types. Additionally, the ability to carry and spread antimicrobial resistance among other Enterococcus spp. has been observed from an analysis of retail meat samples (Jung et al., 2013). The high number of unclassified genera, presented in this study, is possibly indicative of the in

5. Conclusion
The data presented in this study suggests an immense untapped potential for microbiota manipulation in unconventionally grown birds and could reveal useful information for future attempts in standardising the microbiome of industrial poultry. The application of nanoparticles, with careful optimisation, could help uncover a range of unknown bacterial species and their role in the expression of beneficial microbial products. Nanoparticles have rapidly emerged in the food and agricultural industry, and it is of vital importance to understand their gut microbiome interaction, while modifying their properties to our best advantage.

Conflict of interest
We declare that we have no financial and personal relationships with other people or organizations that can inappropriately influence our work, there is no professional or other personal interest of any nature or kind in any product, service and/or company that could be construed as influencing the content of this paper.

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Appendix. Supplementary data
Supplementary data to this article can be found online at https://doi.org/10.1016/j.aninu.2019.06.004.

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