SHORT COMMUNICATION

Identifying bacterial predictors of honey bee health

Giles E. Budge¹,²*, Ian Adams, Richard Thwaites¹, Stéphane Pietravalle¹, Georgia C Drew³,
Gregory D.D. Hurst³, Victoria Tomkies¹, Neil Boonham¹,², Mike Brown⁴

¹ Fera, Sand Hutton, York, YO411LZ.
² Institute for Agri-Food Research and Innovation, Newcastle University, Newcastle upon
Tyne, NE1 7RU.
³ Institute of Integrative Biology, University of Liverpool, Crown Street, Liverpool, L69 7ZB.
⁴ Animal and Plant Health Agency, Sand Hutton, YO41 1LZ.

* Corresponding author: giles.budge@fera.co.uk; giles.budge@ncl.ac.uk
Abstract

Non-targeted approaches are useful tools to identify new or emerging issues in bee health. Here, we utilise next generation sequencing to highlight bacteria associated with healthy and unhealthy honey bee colonies, and then use targeted methods to screen a wider pool of colonies with known health status. Our results provide the first evidence that bacteria from the genus Arsenophonus are associated with poor health in honey bee colonies. We also discovered Lactobacillus and Leuconostoc spp. were associated with healthier honey bee colonies. Our results highlight the importance of understanding how the wider microbial population relates to honey bee colony health.

Keywords

probiotic; symbiont; microbiome
1. Introduction

The economic contribution of insect pollination to crop production (Gallai et al., 2009) and human nutrition security (Ellis et al., 2015) is significant. Managed honey bees are often singled out as a substantial global supplier of pollination services (Kleijn et al., 2015) but are exposed to a range of pressures that contribute to poor health, including parasites (Budge et al., 2015; Higes et al., 2008), pesticides (Henry et al., 2012) and climate change; for review see (Vanbergen and Initiative, 2013).

As pollinators are placed under increasing pressures, the microbiome of bees is emerging as an important and understudied factor in the maintenance of health. Food amended with lactic acid bacteria can protect honey bees against American (Forsgren et al., 2010) and European foulbrood (Vasquez et al., 2012) whilst members of the gut microbiota have putative roles in the metabolism of carbohydrates (Lee et al., 2015). Microbiota of the honey bee may therefore contribute to pathogen defence, nutrition and protection against environmental compounds.

Here we used pyrosequencing of the 16S amplicon to highlight bacteria differentially associated with healthy and unhealthy honey bee colonies, and then developed targeted real-time PCR methods to explore microbial relationships with colony health.
2. Materials and Methods

2.1. Sampling

A recent study collected adult honey bee samples from healthy and unhealthy UK colonies to investigate known pathogens as predictors of poor honey bee colony health (Budge et al., 2015). We identified two case studies within these samples where professional beekeepers managed apiaries experiencing persistently poor colony health as well as apiaries showing consistently good colony health, despite using similar beekeeping practices. Beekeeper A had one healthy apiary (AH; 6 colonies) and two unhealthy apiaries (AU1; 5 colonies and AU2; 6 colonies). Beekeeper B had one healthy apiary (BH; 3 colonies) and one unhealthy apiary (BU; 3 colonies). DNA was extracted from 30 adult honey bees from each colony as described previously (Budge et al., 2015).

2.2. Pyrosequencing 16S amplicons

16S amplicons were produced using composite primers (Hamady et al., 2008) with Multiplex Identifiers (MIDs) from Roche using a different MID tagged reverse primer for each sample (Table S1). The forward primer comprised the Roche 454 Primer B (underlined) and ‘TC’ linker (italics) concatenated to the conserved bacterial primer 27F (bold) (5’-GCCTTGCCAGCCCGCTCAGTCAGCTTGGATCCTGGCAGTCAG-3’). The reverse primer comprised the Roche 454 Primer A (underlined) followed by the 10 nt MID, a ‘CA’ linker (italics) and the conserved bacterial primer 338R (bold) (5’-GCCTCCCTCGCCATCATCA-GCCTCCCTCCTCCCGCTTAGAGT-3’).

16S PCR reactions were set up using Advantage 2 Reagents (Clontech, USA) comprising 5 μL 50x SA buffer, 1 μL Advantage 2 polymerase mix, 0·2 mM dNTPs, 1 μL of template 400 nM forward and reverse primers and 40 μL water. Reactions were carried out in a Biometra T3 thermocycler PCR machine (Biometra, Germany) beginning with 94˚C for 10 min followed by 30 cycles of 95˚C for 30 s (denaturing), 55˚C for 30 s (annealing) and 72˚C for 1 min (extension). PCR products were visualised on a 1% gel and quantified using the Quant-iT
dsDNA BR assay kit (Invitrogen). Amplicons were sequenced on two sixteenths of a plate from a GS-FLX Genome Sequencer (University of Newcastle, Institute of Human Genetics) and sequences analysed using the Ribosomal Database Project (RDP) pyrosequencing pipeline (Cole et al., 2009). Sequences were trimmed and identified based on MID using the initial processing feature and each read assigned to a taxa using the RDP classifier.

2.3. Screening colonies with known health status

Three bacterial species with differential expression between healthy and unhealthy hives were selected for the development of targeted real-time PCR tests following previously published protocols (Budge et al., 2010) (Table S2). Targeted real-time PCR tests were used to rescreen DNA extracts from 129 adult honey bee samples reported previously (Budge et al., 2015). To investigate the relationship between the presence of the newly identified bacteria and honey bee colony health, the square root of the number of combs of adult bees was used as the response variable in a multiple linear regression model with the detection of established parasites (N. apis, N. ceranae, M. plutonius, KBV, DWV, BQCV, SBV, CBPV, APBV, IV, IAPV) and newly associated bacterial species (Arsenophonus, Lactobacillus, Leuconostoc) as potential explanatory variables (GenStat version 17.1).

2.4. Arsenophonus PCR sequencing

To further characterise Arsenophonus spp. detected in A. mellifera adults, we generated sequence from two genes; the house keeping gene fructose-bisphosphate aldolase class II (fbaA) and 16S rRNA for two colony samples using established protocols. FbaA sequences were amplified using the primer pair fbaAF (5′-GCCGCTAAGGTTCTCC) and fbaAR (5′-CCTGAACCACCATGAAAAA; 658 bp amplicon) adapted from a previous study (Duron et al., 2010). 16S rRNA sequences were amplified using established primers (Duron et al., 2008) generating a 804 bp amplicon. Products were purified and Sanger sequenced through both strands using the original primers. Data were used to infer the relatedness of the A. mellifera Arsenophonus strain to others in the genus. Model selection was made
using the best-fit nucleotide substitution test in MEGA6 (Tamura et al., 2013), and maximum
likelihood tree estimated using the Tamura 3-parameter model (Tamura, 1992) for fbaA
sequence, and the Kimura 2-parameter model (Kimura, 1980) for 16S rRNA. The
evolutionary rate differences between sites was modelled using Gamma distribution (fbaA)
or uniform rates (16S rRNA). Accession numbers and references for sequences from the
related species used in phylogenetic reconstruction are provided (Tables S3, S4)
3. Results and Discussion

3.1. Pyrosequencing 16S amplicons

In total, 15,633 16S amplicon sequences (NCBI Bioproject PRJNA315609) were identified by MID and classified with 95% confidence using the RDP webtools. Bacteria from 17 identifiable genera generated at least 1% of the sequence reads in samples from either healthy or unhealthy honey bee colonies (Table 1).

Table 1

Sequences of *Arsenophonus* were more frequently found in adult bee samples from unhealthy apiaries whilst *Lactobacillus* and *Leuconostoc* were more frequently found in healthy apiaries (Table 1). These bacterial genera were selected for further study and real-time PCR primers designed to confirm species presence (Table S2).

3.2. Screening colonies with known health status

PCR-based rescreening of DNA samples from adult honey bees for the remaining three bacterial genera revealed positive results for *Arsenophonus* (62/129), *Lactobacillus* (20/129) and *Leuconostoc* (18/129). The multiple linear regression suggested the established parasite DWV and newly associated bacterial species *Arsenophonus*, *Lactobacillus* and *Leuconostoc* were significant predictors of honey bee colony size (F=20.81; df=4,124; P<0.001). DWV (F=18.68; df=1,124; P<0.001) and *Arsenophonus* (F=9.4; df=1,124; P=0.003) presence were negatively correlated and *Lactobacillus* (F=4.14; df=1,124; P=0.044) and *Leuconostoc* (F=51.01; df=1,124; P<0.001) were positively correlated to the number of combs of bees (Figure 1A).

Figure 1

3.3. *Arsenophonus* PCR sequencing

*Apis mellifera* *Arsenophonus* grouped with *Arsenophonus* strains previously identified in *Colletes* using 16S Sequence (Figure 2A), a result congruent with results from Switzerland (Yañez et al., 2016). FbaA sequences suggested *Apis mellifera* *Arsenophonus* formed a
monophyletic group with *Arsenophonus nasoniae* from the parasitoid wasp (*Nasonia vitripennis*) and *Arsenophonus* isolated from the raspberry aphid (*Aphis idaei*; Figures 1C).
4. Discussion

Our results provide the first evidence that members of the genus *Arsenophonus* are associated with poor health in UK honey bee colonies. In total, 48% of adult bee samples tested positive from eleven counties demonstrating *Arsenophonus* is well distributed geographically, and more common in the UK than Switzerland where only 24% of colonies tested positive (Yañez et al., 2016). Increased abundance of bacteria with 90% sequence identity to *Arsenophonus* has been reported in honey bee colonies suffering from Colony Collapse Disorder (CCD) in the United States, indicating a potential association with poor bee health (Cornman et al., 2012). There are two competing and equally significant hypotheses for the correlation between *Arsenophonus* presence and the poor health of honey bee colonies. Firstly, *Arsenophonus* could increase host susceptibility to infection. This might occur, for instance, if the symbiont modulated host immune pathways are affected to reduce pathogen clearance. Alternatively, *Arsenophonus* may protect its host against parasites, and thus reaches high prevalence in areas where parasite pressure is high. *Arsenophonus* has been associated with foraging honey bees in Israel (Aizenberg-Gershtein et al., 2013), Switzerland (Babendreier et al., 2007) and The United States (Corby-Harris et al., 2014) and was associated with hive debris from the Czech republic (Hubert et al., 2015), so whilst we do not know which of our hypotheses is correct, elucidation of the association is of clear importance to international apiculture and merits future experimental studies.

We also report the novel finding that lactic acid bacteria (LAB) from the genera *Lactobacillus* and *Leuconostoc* were predictors of increased colony size in UK honey bee colonies. *Leuconostoc* spp. have rarely been associated with aculeate pollinators, the only previous reports being presence in fresh pollen collected by foraging honey bees in Algeria (Belhadj et al., 2010) and a finding in the gut of *Bombus terrestris* in Belgium (Praet et al., 2015). *Lactobacillus* is better studied, becoming associated with adult bees soon after eclosure (Vasquez et al., 2012) and thought to be important to honey production (Olofsson and
Vasquez, 2008) and the maturation of pollen (Vasquez and Olofsson, 2009). LABs have long been associated with good health in humans and although they have recently been shown to inhibit bacterial honey bee pathogens (Forsgren et al., 2010; Vasquez et al., 2012) our data are the first to link their presence with good colony health. Several commercial feeds contain blends of LAB (including *Lactobacillus*) to offer the promise of improved honey bee colony vigour, however none of these products are known to contain *Leuconostoc* spp.. Future experiments should determine whether the inclusion of *Leuconostoc* spp. could improve the health of honey bee colonies as part of a novel probiotic.

Our results contribute to the growing body of evidence that the honey bee microbiota, outwith known pathogens, may offer an important contribution to honey bee colony health. Non-targeted sequencing methods are a useful tool to highlight previously unknown microbes and other genera, such as *Microbacterium*, *Proteus* and *Staphylococcus*, represent additional possible candidates for further study.
Acknowledgments

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References


causes honeybee colony collapse. Environmental Microbiology 10, 2659-2669.


Table and figure legends

Table 1  Frequency of 16S amplicon sequences detected in adult honey bee samples for all 17 identifiable genera with greater than 1% read abundance in either healthy or unhealthy groups.

Figure 1  Estimated number of combs of adult bees as predicted by presence or absence of deformed wing virus (DWV), Arsenophonus, Lactobacillus and Leuconostoc using a multiple linear regression (A). Error bars represent 95% CI. Maximum likelihood inference of the relatedness of Arsenophonus spp. isolated from Apis mellifera to other Arsenophonus strains using sequence from 16S rRNA (B) and fbaA (C). Branch length denotes the number of substitutions per site and bootstrap values from 1000 replications are shown at nodes. Strains that have not been formally taxonomised are labelled following their host species.