

Microbiome and infectivity studies reveal complex polyspecies tree disease in Acute Oak Decline

Denman, Sandra; Doonan, James; Ransom-Jones, Emma; Broberg, Martin; Plummer, Sarah; Kirk, Susan A.; Scarlett, Kelly ; Griffiths, Andrew; Kaczmarek, Maciej; Foster, Jack ; Peace, Andrew; Golyshin, Peter; Hassard, Francis; Brown, Nathan ; Kenny, John G.; McDonald, James

The ISME Journal

DOI:
[10.1038/ismej.2017.170](https://doi.org/10.1038/ismej.2017.170)

Published: 01/01/2018

Peer reviewed version

[Cyswllt i'r cyhoeddiad / Link to publication](#)

Dyfyniad o'r fersiwn a gyhoeddwyd / Citation for published version (APA):

Denman, S., Doonan, J., Ransom-Jones, E., Broberg, M., Plummer, S., Kirk, S. A., Scarlett, K., Griffiths, A., Kaczmarek, M., Foster, J., Peace, A., Golyshin, P., Hassard, F., Brown, N., Kenny, J. G., & McDonald, J. (2018). Microbiome and infectivity studies reveal complex polyspecies tree disease in Acute Oak Decline. *The ISME Journal*, 12, 386-399.
<https://doi.org/10.1038/ismej.2017.170>

Hawliau Cyffredinol / General rights

Copyright and moral rights for the publications made accessible in the public portal are retained by the authors and/or other copyright owners and it is a condition of accessing publications that users recognise and abide by the legal requirements associated with these rights.

- Users may download and print one copy of any publication from the public portal for the purpose of private study or research.
- You may not further distribute the material or use it for any profit-making activity or commercial gain
- You may freely distribute the URL identifying the publication in the public portal ?

Take down policy

If you believe that this document breaches copyright please contact us providing details, and we will remove access to the work immediately and investigate your claim.

Title: Microbiome and infectivity studies reveal complex polyspecies tree disease in Acute Oak Decline

Authors: Sandra Denman^{1*†}, James Doonan^{2*}, Emma Ransom-Jones², Martin Broberg², Sarah Plummer¹, Susan Kirk¹, Kelly Scarlett¹, Andrew R. Griffiths^{1,2}, Maciej Kaczmarek^{1,2}, Jack Forster¹, Andrew Peace³, Peter N. Golyshin², Francis Hassard⁴, Nathan Brown⁵, John G. Kenny⁶, James E. McDonald^{2†}.

*These authors contributed equally to this work; † joint corresponding authors.

Affiliations:

¹Forest Research, Centre for Forestry and Climate Change, Alice Holt Lodge, Farnham, Surrey GU10 4LH, UK.

²School of Biological Sciences, Bangor University, Deiniol Road, Bangor, Gwynedd, LL57 2UW, UK.

³Craigernie Drive, Peebles, Scottish Borders, EH45 9HN, UK.

⁴School of Ocean Sciences, Bangor University, Bangor, Anglesey, LL59 5AB, UK.

⁵Department of Computational and Systems Biology, Rothamsted Research, West Common, Harpenden, Hertfordshire, AL5 2JQ, UK.

⁶Centre for Genomic Research, Institute of Integrative Biology, University of Liverpool, Liverpool, L69 7ZB, UK.

*Correspondence to: James McDonald, j.mcdonald@bangor.ac.uk; Sandra Denman, Sandra.Denman@forestry.gsi.gov.uk

The authors declare no conflicts of interest.

22 **Subject category: Microbe-microbe and microbe-host interactions**

23 **Abstract**

24 Decline-diseases are complex and becoming increasingly problematic to tree health
25 globally. Acute Oak Decline (AOD) is characterised by necrotic stem lesions and
26 galleries of the bark-boring beetle, *Agrilus biguttatus*, and represents a serious threat to
27 oak. Although multiple novel bacterial species and *Agrilus* galleries are associated with
28 AOD lesions, the causative agent(s) are unknown. The AOD pathosystem therefore
29 provides an ideal model for a systems-based research approach to address our hypothesis
30 that AOD lesions are caused by a polymicrobial complex. Here we show that three
31 bacterial species, *Brenneria goodwinii*, *Gibbsiella quercinecans* and *Rahnella victoriana*,
32 are consistently abundant in the lesion microbiome, and possess virulence genes used by
33 canonical phytopathogens that are expressed in AOD lesions. Individual and polyspecies
34 inoculations on oak logs and trees demonstrated that *B. goodwinii* and *G. quercinecans*
35 cause tissue necrosis, and in combination with *A. biguttatus* produce the diagnostic
36 symptoms of AOD. We have proved a polybacterial cause of AOD lesions, providing
37 new insights into polymicrobial interactions and tree disease. This work presents a novel
38 conceptual and methodological template for adapting Koch's postulates to address the
39 role of microbial communities in disease.

Introduction

Trees are essential to landscape function and aesthetics, supporting diverse ecologies (Rackham, 2008) and providing key ecosystem services (Boyd *et al.*, 2013). However, significant areas of forest have been lost due to increasing outbreaks of disease and pest attack, and tree health is a current global concern (Cohen *et al.*, 2016). Tree diseases, including decline-diseases, are rising in profile due to an increased risk of introduction and spread through international plant trade, and amplification effects of current and future climate change (Millar and Stephenson, 2015; McDowell *et al.*, 2011; Oliva *et al.*, 2014). Emerging evidence suggests that complex biotic interactions, including polymicrobial and insect activity affect disease occurrence and severity (Adams *et al.*, 2013; Buonauro *et al.*, 2015; Lamichhane and Venturi, 2015), yet little progress has been made in applying the latest advances in sequencing and culture-based methodologies to characterize pathosystems in trees. The prevailing paradigm of infection biology contends that one organism causes one disease (proved using Koch's postulates). In contrast, there is increasing recognition of the importance of polymicrobial interactions in human disease, following developments in sequencing technologies that allow microbiome-wide association studies to identify the role of microbial communities in disease (Gilbert *et al.*, 2016). In the medical field this is leading to adaptations of Koch's postulates to include complex interactions between the environment, host, and microbial communities (Fredricks and Relman, 1996; Falkow, 1988; Hill, 1965). However, progress in characterising polyspecies interactions in plant disease has been limited, although there is clearly a need for contemporary approaches to investigating complex tree diseases (Lamichhane and Venturi, 2015). Analysis of such complex biotic

63 interactions requires an integrated, systems approach, particularly in the case of decline-
64 diseases where both complex abiotic and biotic interactions underpin disease
65 development.

66 Decline-diseases, first formally described as a specific disease in the United States
67 (Sinclair, 1965; Manion, 1981) but well documented elsewhere (Delatour, 1983; Thomas,
68 2008; Sinclair and Lyon, 2005), are of global concern (Pautasso *et al.*, 2015). Unlike
69 most common tree diseases, decline-diseases are not caused by single primary pests or
70 pathogens; instead they are complex syndromes, involving the sequential, combined and
71 cumulative effects of (often secondary) biotic and abiotic agents (Brown *et al.*, 2016;
72 Sallé *et al.*, 2014; Thomas, 2008; Manion, 1981). Currently the UK is facing an episode
73 of Acute Oak Decline (AOD), which occurs widely in southern and midland England,
74 extending into Wales (Brown *et al.*, 2016; Denman *et al.*, 2016) and represents a
75 significant threat to oak, particularly native species *Quercus robur* and *Q. petraea*
76 (Denman *et al.*, 2014). First recognized in Britain in the 1980s (Denman and Webber,
77 2009), similar declines have occurred in continental Europe (Gibbs and Greig, 1997;
78 Hartmann *et al.*, 1989; Biosca *et al.*, 2003; Vansteenkiste *et al.*, 2004). AOD-affected
79 trees show discrete, weeping stem patches (stem bleeds), signifying areas of necrosis and
80 fluid filled cavities in the underlying inner bark (Denman *et al.*, 2014) (see Figure 1a-c),
81 which disrupt vascular flow of nutrients and water essential to tree survival
82 (Vansteenkiste *et al.*, 2004). Larval galleries of the buprestid beetle *Agilus biguttatus* are
83 found in conjunction with lesions (Brown *et al.*, 2015, 2017) (Figure 1d), and can also
84 impact tree condition by girdling the tree when colonization is intense, leading to tree

death (Sallé *et al.*, 2014). AOD typically affects mature oaks, but has also been reported in young trees (Brown *et al.*, 2016).

First steps to determine the causes of stem bleeds led to the isolation of several novel bacterial species from AOD lesions, with three novel species, *Gibbsiella quercinecans* (*Enterobacteriaceae*) (Brady *et al.*, 2010), *Brenneria goodwinii* (*Pectobacteriaceae*) (Adeolu *et al.*, 2016; Denman *et al.*, 2012), *Rahnella victoriana* (*Yersiniaceae*) (Adeolu *et al.*, 2016; Brady *et al.*, 2014), and an un-named *Pseudomonas* (Denman *et al.*, 2016; Sapp *et al.*, 2016) consistently isolated. Some of these species, e.g. *G. quercinecans* as *Serratia* sp. (Biosca *et al.*, 2003; Poza-Carrión *et al.*, 2008) on *Quercus pyrenacia* and *Q. ilex* are implicated as causative agents of stem bleeding on other oak species in Europe (Biosca *et al.*, 2003; Poza- Carrión *et al.*, 2003; Denman *et al.*, 2016). There was less consistent isolation of various other bacterial species, for example *Lonsdalea quercina* ssp. *britannica* (Brady *et al.*, 2012), which is closely related to *Lonsdalea quercina* ssp. *quercina*, the causative agent of acorn gummosis on *Q. agrifolia* and *Q. wislizenii* in America (Hildebrand and Schroth, 1967), and several other novel species that are in the process of being formally described. Although correlation of certain bacterial species with AOD symptomology has been observed, empirical evidence on the causative agent(s) of AOD lesions is lacking and remains a barrier to developing informed management strategies for this disease. In the absence of a single putative primary pathogen as the causal agent of AOD lesions, we hypothesised that a polymicrobial complex is responsible for AOD lesion formation. However, demonstrating causation by a polymicrobial complex with associated insect activity on mature oak trees or logs is

challenging in the context of fulfilling Koch's postulates and requires a multi-faceted methodological and conceptual approach.

Here, we applied a systems-level approach to determine cause(s) of necrosis in AOD using microbial isolation and culture, phenotypic tests, genomic analyses of *G. quercinecans*, *B. goodwinii*, and *R. victoriana*, metagenome and metatranscriptome analysis of healthy and diseased trees, and inoculation tests comprising both polybacterial inoculations and the addition of live *A. biguttatus* eggs to recreate the symptoms of AOD. This combined sequencing and cultivation-based approach provides a contemporary adaptation of Koch's postulates to address the biotic components of a complex decline-disease, and represents a conceptual model for future analyses of polymicrobial infections in trees and other systems.

Methods

Isolation of bacteria from healthy and diseased oak trees

In the search for putative causal agents of stem lesions we wanted to determine the veracity of differences in occurrence and composition of bacterial communities in lesions and visually healthy trees, as the bacterial microbiome has previously been identified as the likely causal agent of stem lesions (Denman *et al.*, 2016). Conventional isolation and culture and microbiome analyses were used. Samples were acquired through citizen science reports (CSR), as well as structured studies (Denman *et al.*, 2016; Sapp *et al.*, 2016) ensuring the broadest possible coverage of AOD sites (Supplementary Table S1 and Supplementary Table S2). Eighteen CSR sites were sampled (Supplementary Figure

S1 and Supplementary Table S1), together with those from structured studies (Denman *et al.*, 2016; Sapp *et al.*, 2016), as well as five sites that had no history of AOD (N-AOD). Trees were sampled by forest pathologists who made site visits following the enquiry. In total, isolations from 66 trees were analysed. Destructive sampling of trees, by removing panels of diseased oak tissue, as well as healthy oak tissue from apparently healthy trees on the same sites, was carried out where permitted, and isolations were made using PYGA medium as described in Denman *et al.* (2014; 2016) and Sapp *et al.* (2016). Owing to the nature of the CSR studies, the number of tissue pieces plated out was variable, dependent upon the sample. Bacterial colonies emerging from chips of tissue were purified using standard streak-plating techniques; single visually representative colonies were selected, cultivated in Luria Bertani broth (LB) and identified with PCR and DNA amplicon sequencing as described in (Denman *et al.*, 2016).

Statistical analysis of isolation study datasets

Bacterial yield between healthy and diseased trees was tested by fitting a generalized linear mixed effects model with logit link function and binomial error distribution. Fixed effects were fitted for tree health and tissue position and random effects fitted for sites and trees within sites. Over-dispersion in the model was taken into account by including an additional dispersion parameter.

Differences in bacterial communities were analyzed by detrended correspondence analysis, down weighting bacterial species occurring in less than 5% of tissue combinations. Monte Carlo permutation tests were used to test for significant differences

in bacterial communities between healthy and diseased trees and to test the effect of tissue position within the tree. Finally, Jaccard's similarity index was used to identify any significant associations between bacteria across the 66 trees of the study.

Genome sequencing of bacterial strains isolated from Acute Oak Decline affected trees

Maintenance of bacterial strains used in genome analyses

Gibbsiella quercinecans FRB97 (T) (Brady *et al.*, 2010), *Brenneria goodwinii* FRB141 (T) (Denman *et al.*, 2012), and *Rahnella victoriana* BRK18a (T) (Brady *et al.*, 2014) were isolated by Forest Research in the CSR studies, from oak trees affected by AOD. Strains were stored in glycerol stocks at -80°C and maintained on nutrient agar (Oxoid) at 20°C.

DNA preparation and genome sequencing on Pacific Biosciences RSII sequencing platform

The whole genomes of *G. quercinecans* FRB97, *B. goodwinii* FRB141 and *R. victoriana* BRK18a were sequenced using the Single Molecule Real-Time (SMRT) technology of the Pacific Biosciences RSII platform. A single colony of each isolate was sampled from nutrient agar (Oxoid) and used as inoculum for liquid culture, which was grown overnight in nutrient broth (Oxoid) at 28°C, and shaken at 150 rpm. Total genomic DNA was isolated using the Gentra Puregene Yeast/Bact. kit (Qiagen) and quantified using a

Qubit fluorometer (Life Technologies, Paisley, UK). DNA integrity was assessed using 1% agarose gel electrophoresis. *G. quercinecans* and *B. goodwinii* DNA libraries were prepared using 10 µg of genomic DNA and sequenced by DUGSIM at Duke University, NC, USA, using P4/C2 chemistry and six SMRT cells per isolate. The *R. victoriana* DNA library was prepared and sequenced by the Centre for Genomic Research, University of Liverpool, UK using P6/C4 chemistry and one SMRT cell. All sequence data described in this study is available under BioProject PRJNA323828, (Supplementary Table S3). Genome assembly and annotation is described in Supplementary Methods.

Metagenome sequencing of diseased and healthy oak trees

Collection of samples for metagenome sequencing

Tissue samples were collected from Runs Wood, Ross-on-Wye, and two sites in Attingham park (Supplementary Table S2). More than half of the trees sampled for metagenome analysis were also sampled in the isolation study. A panel comprising all layers of stem tissue (outerbark, innerbark, sapwood and heartwood) was removed from the visible bleed area on each diseased tree, or from stem areas at similar height on healthy trees, according to previously described methods (Denman *et al.*, 2014, 2016; Sapp *et al.*, 2016). Samples were immediately flash frozen on dry ice and stored at -80°C prior to processing.

Metagenome assembly, annotation and mapping to the genomes of *G. quercinecans* FRB97 (T), *B. goodwinii* FRB141 (T) and *R. victoriana* BRK18a (T)

194 Metagenomic reads were assembled using RAY-meta v2.3.1 (Boisvert *et al.*, 2012) using
195 default parameters. Assemblies were annotated using Prokka v1.11 (Seemann, 2014).
196 Translated annotations were aligned against the translated protein sequences of *G.*
197 *quercinecans* FRB97 (T), *B. goodwinii* FRB141 (T), *R. victoriana* BRK18a (T), and two
198 control genomes *Pectobacterium carotovorum* ssp. *carotovorum* PC1 and *Paenibacillus*
199 *polymyxa* SC2, using BLASTx v2.2.26 (Altschul *et al.*, 1990). Metagenome sequences
200 with greater than 97% homology for at least 50 amino acids to proteins identified in *G.*
201 *quercinecans*, *B. goodwinii* and *R. victoriana* were considered a match (Supplementary
202 Table S4). Control genomes were selected to measure the stringency of alignment. *P.*
203 *carotovorum* ssp. *carotovorum* is a canonical plant pathogen, and a member of the soft-
204 rot *Enterobacteriaceae* (SRE) (N.B. many members have recently been reclassified into
205 novel families (Adeolu *et al.*, 2016; Charkowski *et al.*, 2012)), which had been identified
206 sporadically and at low relative abundance from metagenomic taxonomic surveys within
207 this study (Supplementary Table S5). However, *P. carotovorum* ssp. *carotovorum* had
208 not previously been isolated from necrotic lesions on Acute Oak Decline affected trees.
209 Therefore, it was proposed that coding domain alignments to *P. carotovorum* ssp.
210 *carotovorum* align to conserved genes which are present in many *Enterobacteriaceae*,
211 and their low relative abundance does not signify their presence but is an artefact of the
212 alignment process. Resultant Circos plots agree with this proposal as conserved genes are
213 frequently found in the metagenomic alignment against the *P. carotovorum* ssp.
214 *carotovorum* PC1 genome, which are likely to align to other *Enterobacteriaceae*. Within
215 *G. quercinecans*, *B. goodwinii* and *R. victoriana* there is strong homology to conserved
216 and variant genes. *P. polymyxa* SC2 was identified at low relative abundance in healthy

and diseased metagenomic samples (Supplementary Table S5), therefore it was selected to test stringency of the metagenomic alignment and to measure its relative abundance in healthy and diseased microbiomes using an alternative method. Homologous bacterial protein identities and the workflow used for metagenomic analysis is available from GitHub: (https://github.com/clydeandforth/multi_omics_study.git).

Taxonomic classification of metagenome sequences

To compare the taxonomic composition of the oak microbiome, raw sequence reads were taxonomically labelled using Kraken v0.10.5 beta (Wood and Salzberg, 2014) and One Codex (Minot *et al.*, 2015) (Figure 2a). Taxonomic labelling using Kraken was performed on the standard RefSeq genome database supported by Kraken, with the addition of the genomes of *B. goodwinii* FRB141 (T), *R. victoriana* BRK18a (T), *G. quercinecans* FRB97 (T), and *Lonsdalea quercina* ssp. *quercina* ATCC 29281 (Figure 2d and Supplementary Table S5). Taxonomic analysis using One Codex was conducted against the One Codex March '16 Preview database with the addition of the genomes of *G. quercinecans*, *B. goodwinii* and *R. victoriana*.

Functional annotation of metagenome sequences

Metagenome datasets derived from samples AT2, AT3, AT4, AT5, AT6, ROW1, ROW2 and ROW3 were analysed via MG-RAST (Meyer *et al.*, 2008) using Hierarchical Classification against the Subsystems database with an *E*-value cut-off of 1e-5, a minimum percentage identity cut-off of 80%, and a minimum alignment length of 50.

Descriptions of the taxonomic and functional composition of the metagenomes derived from MG-RAST were comparable with those derived from analysis of the same datasets using One Codex and Kraken, and further validated by mapping of metagenome reads against the finished genomes of *B. goodwinii*, *G. quercinecans* and *R. victoriana*.

Statistical analysis of metagenome datasets

Statistical analyses were performed using Primer v7 (Clarke and Gorley, 2015) with PERMANOVA+ add on to explore relationships between community changes (Figure 2b). One Codex metagenome data was log (N+1) transformed, to downweight the most abundant genera. Next, dissimilarities were calculated with the S17 Bray-Curtis similarity coefficient. A principal coordinate ordination analysis was performed by plotting the inter-point dissimilarity values for each factor (site and disease status), the variation in community composition was plotted as the first two axes (preserving actual dissimilarities) (Gower, 1966). A correlation was performed between each taxon and each community coordinate. Correlations with each component were deemed significant ($R^2 > 0.5$) and a vector biplot was overlaid to visualise the strength of the correlation. A Welch's *t*-test was performed to test significance of differences between key taxa (identified above) healthy and diseased trees (pooled abundances for each factor). Resultant *p*-values from Welch's *t*-test are overlaid on correlation biplot with significance at >95% ($p < 0.05$) deemed significant. Comparative functional analysis of MG-RAST (Meyer *et al.*, 2008) annotated, metagenome data was performed using Stamp v2.1.3 (Parks *et al.*, 2014). Statistically significant functional differences between diseased and healthy communities were calculated using a G-test with Yates' correction. The

Newcombe-Wilson test was performed to calculate confidence intervals between two binomial population proportions (Brown and Li, 2005).

Metatranscriptome sequencing of AOD diseased oak trees

Collection of samples for metatranscriptome sequencing and RNA extraction

For RNA sampling, two separate lesions (samples AT11 and AT12) from a single tree were analysed in June 2013. Swabs of the lesion fluid were collected in addition to tissue from the active margins of the lesion and immediately frozen in liquid nitrogen, and transported back to the laboratory in a vessel containing liquid nitrogen. Samples were stored at -80°C prior to processing. Before RNA extraction, samples in liquid nitrogen were ground with a pestle in a mortar to homogenise the tissue. RNA was extracted from 2g of tissue using the PowerSoil Total RNA Isolation Kit (MoBio) according to manufacturer's instructions. RNA was quantified using a Qubit fluorometer (Thermo Fisher) and quality assessed using the Bioanalyzer 2100 (Agilent).

Metatranscriptome sequencing

Sequencing libraries were prepared from samples of total RNA using the strand-specific ScriptSeq preparation kit (Illumina), and sequenced using 2x100bp paired-end sequencing on the Illumina HiSeq platform. Reads were trimmed using first Cutadapt 1.2.1 (Martin, 2011) and additionally Sickle 1.2.00 (Joshi and Fass, 2011). Due to low

RNA yields from the lesion samples, total RNA was sequenced and rRNA sequence reads were subsequently depleted *in silico* prior to mRNA transcript analysis.

Metatranscriptome assembly and functional annotation

Two *in silico* rRNA depleted metatranscriptome libraries were aligned to the *Gibbsiella quercinecans* FRB97 (T), *Brenneria goodwinii* FRB141 (T), *Rahnella victoriana* BRK18a (T), and control genomes (*P. carotovorum* ssp. *carotovorum* PC1 and *P. polymyxa* SC2) (Supplementary Figure S2 and S3) with Bowtie2 v2.2.4 (Langmead and Salzberg, 2012), using local mode to maximise alignment score. Aligned reads were converted from Sequence Alignment/Map (SAM) to Binary Sequence Alignment/Map (BAM) format and indexed using SAMtools v1.2 (Li *et al.*, 2009). To avoid false positives in the detection of gene expression, a gene was considered as being expressed if 3 or more transcripts were aligned and the combined coverage from both libraries represented more than 20% of the gene, (adapted from (Versluis *et al.*, 2015) (Figure 2a-c and Supplementary Figure S4). A custom Perl script was designed to extract transcript alignments, and is available from GitHub: (https://github.com/clydeandforth/multi_omics_study.git). Aligned transcripts were visualised in Artemis (Carver *et al.*, 2012).

Log inoculations

Pathogenicity tests were set up to reproduce lesions characteristic of AOD under controlled conditions. The following hypotheses were tested: 1) Key species consistently

isolated from AOD symptomatic oak can cause necrosis of oak stem tissue, 2) Combinations of key bacterial species cause more severe tissue necrosis (reflected in larger lesions), than individual species alone, 3) The interaction between *A. biguttatus* larvae (derived from eggs) and bacteria, leads to the development of AOD symptoms.

Four experiments were carried out over three consecutive years as testing could only be done annually when beetle eggs were available. Three trials were carried out on oak logs in growth chambers, and the fourth trial was set out in the field, where stems of young plantation-oak (25 years old), were used instead of logs.

Growth Chamber Log trials

Logs used in the trials were obtained from freshly felled *Q. robur* trees, with diameter at 1.3m (DBH) = 16 - 20 cm. The trees were located in the Straits Enclosure of the Alice Holt Forest, Hampshire, England, UK, and logs were transported to the laboratory after felling where they were prepared for inoculation. Logs measuring 40 cm (mini-logs), or 130 cm (long logs) in length had the uppermost cut-surface sealed with isoflex liquid rubber (Ronseal Ltd, Sheffield) to prevent desiccation. Logs were placed, lower cut surface down, in saucers containing water. Logs were inoculated in August or September, and incubated at 25°C in the growth chamber with a 12h photoperiod. Mini-logs were used for all treatments involving *A. biguttatus* eggs, in each case only one bacterial + eggs treatment type per mini-log was tested to guard against cross-contamination through larval spread (Supplementary Table S2). All the bacterial treatments without eggs (either single species or combinations of species) were placed on the same long log, with

inoculation points marked out along its length. Eggs of *A. biguttatus* were produced in Forest Research's laboratories at Alice Holt (Reed, 2016). In a single experiment (2014 NW; Supplementary Table S2), non-wound inoculations were carried out; the remaining two growth chamber trials and the field trial were inoculated using shallow wounds to the outer surface of the innerbark, made by a 10 mm cork-borer. Half a loop of inoculum scooped from 24h-old inoculum plates using disposable plastic loops was inserted into the wound and rubbed to dislodge the bacterial cells around the wound surface. The outerbark plug was replaced on the inoculation point, wounds covered with parafilm and damp cotton wool, and sealed with duct tape. After 4 months incubation, experiments were terminated, outer bark stripped from inoculation points to expose lesions, which were hand traced onto tracing paper, and back isolations were cultured onto peptone yeast glucose agar. Mass bacterial colonies that developed were tested for the presence of *B. goodwinii*, *G. quercinecans*, *Rahnella* and *Lonsdalea* using a multiplex Taqman qPCR assay. Lesion areas were calculated using ASSESS V2 (APS, Minnesota).

Statistical analyses

Log lesion areas

The four pathogenicity trials were used to establish the impact of different bacterial species tested individually, in combination (polybacterial inoculations), and with or without the addition of *A. biguttatus* eggs (Supplementary Table S2) on tissue necrosis (assessed by the size of the lesion area associated with each inoculation point).

Lesion area data were refined in a hierarchical fashion, such that the following data sets were used for lesion area analysis:

- 348 1) Non-contaminated samples fulfilling Koch's postulates, with galleries present (for
349 *A. biguttatus* samples indicating that eggs had hatched)
- 350 2) Non-contaminated samples fulfilling Koch's postulates, with or without galleries
351 present (for *A. biguttatus* samples)
- 352 3) Non-contaminated samples
- 353 Further information on statistical analyses of log lesion areas are described in
354 Supplementary Methods.
- 355
- 356
- 357

Results

Isolation study

In the isolation study, analysis of 38 diseased trees from 23 sites, plus 13 healthy trees in 11 of these sites, and 15 healthy trees from five sites with no history of AOD, identified 159 bacterial taxa. Higher yields of bacteria were obtained from lesion margin tissue of symptomatic trees compared with healthy tree tissue ($F_{1,28}$, $p<0.001$), and the lesion margin bacterial profile was significantly different to healthy tissues (Monte-Carlo permutation test of 1000 permutations, $p<0.001$) (Supplementary Figure S5 and Supplementary Table S6). Key genera isolated included *Pseudomonas* (comprising multiple taxa and occurring in healthy as well as diseased trees); but *Gibbsiella*, *Brenneria* and *Rahnella* dominated lesion margins (Supplementary Table S6). *G. quercinecans* occurred on all disease sites, and was isolated from 83% of diseased and 4% of healthy trees sampled, comprising 17% of total diseased samples, and $<0.1\%$ of total healthy samples. *B. goodwinii* was more difficult to isolate, but was obtained from 15 sites, accounting for 16% of total diseased and $<0.1\%$ of total healthy samples. Members of the genus *Rahnella* were obtained from 65% of diseased sites, but *R. victoriana* was isolated only from diseased trees (37%), on 9 of the sites. *L. quercina* ssp. *britannica* (Brady *et al.*, 2012) was isolated sporadically on four sites, and a *Pseudomonas* species (*P. fulva*-like) not yet formally identified, was isolated at eight sites from diseased trees only (Supplementary Table S6). There was a significant co-occurrence of *G. quercinecans* and *B. goodwinii* in diseased tissue ($J=0.56$, $p<0.001$), but neither was isolated from trees on sites with no history of AOD.

Oak microbiome analysis

Taxonomic analysis of the metagenome using unassembled metagenome sequence reads against the One Codex (Minot *et al.*, 2015) March '16 Preview database (with the addition of the genomes of *G. quercinecans*, *B. goodwinii* and *R. victoriana*) also revealed a shift in microbiome composition between healthy stem tissue and AOD lesions (Figures 2a and 2b). *Periglandula*, *Burkholderia*, *Streptomyces*, *Bacillus* and *Auriemonas* were the most abundant genera in healthy trees, whereas *Brenneria* dominated diseased tissue (mean read abundance, 37.5%) (Figure 2a). The mean read abundance of *Gibbsiella* (0.9%) and *Rahnella* (3.7%) was also greater in diseased tissue when compared with healthy tissue (both 0.1%). *Pseudomonas*, a diverse genus comprising both endophytic and phytopathogenic bacteria (Vinatzer *et al.*, 2014), had similar mean abundances in both diseased (4.8%) and healthy trees (3.3%). Correlation coefficients and Welch's unequal variances *t*-tests revealed that *Streptomyces* ($t_{\text{Welch's}} = 49.7$, $p=0.004$) and *Periglandula* ($t_{\text{Welch's}} = 821.8$, $p<0.001$) were significantly associated with healthy trees, whereas *Brenneria* ($t_{\text{Welch's}} = 12.4$, $p=0.006$) and *Gibbsiella* ($t_{\text{Welch's}} = 4.7$, $p=0.05$) were strongly correlated with the lesions of diseased trees (Figure 2b and Supplementary Table S7). *Pseudomonas* and *Rahnella* were not strongly correlated with either health state (Supplementary Table S7).

To identify the most abundant species in the lesion microbiome, raw metagenome reads were aligned using Kraken (Wood and Salzberg, 2014) against reference genome databases (see methods), revealing seventeen genomes that were commonly detected in diseased tissue (Figure 2d and Supplementary Table S7). *B. goodwinii*, *G. quercinecans* and *R. victoriana* were detected in the lesion metagenome of all trees with AOD. Overall,

B. goodwinii was the most abundant genome in the lesion microbiome (range, 0.3-49% of metagenome sequence reads; mean, 12%), but was also detected in much lower proportions in healthy trees (0.01% in all samples). *R. victoriana* (range, 0.01-15%; mean, 2.1%) and *G. quercinecans* (range, 0.02-0.8%; mean, 0.3%) were the second and fourth most abundant genomes in the lesions of diseased trees, respectively. Functional metagenome analysis (Figure 2c and Supplementary Figure S6) revealed that genes associated with carbohydrate metabolism, membrane transport, and virulence, defence and disease, are key features of the lesion microbiome, suggesting that many of these functions are encoded in the genome of *B. goodwinii*, *G. quercinecans* and *R. victoriana*.

Genome analysis of AOD-associated bacteria

Whole genome sequencing of *B. goodwinii* FRB141, *G. quercinecans* FRB97, and *R. victoriana* BRK18a (Supplementary Table S8) revealed that they are phylogenetically related to opportunistic phytopathogens belonging to the SRE (Charkowski *et al.*, 2012; Adeolu *et al.*, 2016) and possess a catalogue of virulence genes associated with canonical phytopathogens (Supplementary Table S9). The SRE alternate their lifestyle from benign commensals to brute force necrotrophic-pathogens, which macerate cell wall polysaccharides by releasing plant cell wall degrading enzymes (PCWDEs) (Toth *et al.*, 2006). A genome-wide search for PCWDEs in *G. quercinecans*, *B. goodwinii* and *R. victoriana* revealed the presence of pectinases, cellulases and tannases (Figure 3b, Supplementary Table S9) whose activity has been confirmed phenotypically (Supplementary Figure S7). Furthermore, *G. quercinecans* and *R. victoriana* possess a type II secretion system, an operon which releases most PCWDEs and is therefore the

central virulence facilitator of necrotrophic plant pathogens. *B. goodwinii* encodes a type III secretion system, the principal virulence factor of established hemibiotrophic pathogens such as *Pseudomonas syringae*, which use the operon to evade immune surveillance, allowing bacteria to increase their numbers before releasing necrotic enzymes as nutrients are depleted (Tampakaki *et al.*, 2010).

To address the role of *B. goodwinii*, *G. quercinecans*, and *R. victoriana* in the aetiology of AOD, we aligned metagenome sequences and transcripts recovered from necrotic lesions of AOD-affected trees against their genomes (Figure 3a). Alignment of six AOD lesion metagenomes, revealed an average of 2225 homologous proteins in *B. goodwinii* FRB141, 858 in *G. quercinecans* FRB97 and 396 in *R. victoriana* BRK18a (Supplementary Table S4). Furthermore, annotated genes from the assembled metagenome of a healthy oak revealed only two homologous proteins in *B. goodwinii* FRB141, one in *G. quercinecans* FRB97, and two in *R. victoriana* BRK18a. Lesion transcripts were aligned to coding regions within *B. goodwinii* FRB141, *G. quercinecans* FRB97, and *R. victoriana* BRK18a; this revealed that the transcripts aligned significantly to virulence genes including PCWDEs, secretion system machinery, and regulators of PCWDEs (Figure 3c and Supplementary Figure S4). *G. quercinecans* FRB97 expressed the pectic enzymes, polygalacturonase and rhamnogalacturonate lyase, and β -glucosidase (cellulase), pectate exo-lyase, oligogalacturonide lyase (which cleaves polygalacturonic acid, the by-product of pectin lyase) (Moran *et al.*, 1968). *B. goodwinii* FRB141 expressed phosphocellobiose, β -galactosidase and several type III secretion system effectors, *R. victoriana* BRK18a expressed many general secretory pathway (GSP) genes, a β -glucosidase, a tannase and carbohydrate esterase enzyme (Yao *et al.*, 2013). Key

regulators of pectinolysis were expressed in all three bacteria including *kdgR*, *phoP*, *pecT*, *rsmA/rsmB*, and *gacA* (Figure 3a).

Log lesion areas

Figures 1e-l show lesions caused by bacteria with or without *A. biguttatus* inoculated into logs in pathogenicity tests. Notable results for lesion area analysis are presented in Figures 1m and 1n (non-contaminated samples fulfilling Koch's postulates, with galleries present for *A. biguttatus* samples). In bacteria only inoculations, lesions significantly bigger than wound controls (Figure 1m) were made by the combination of bacterial species *B. goodwinii* + *G. quercinecans* ($t_{dunnetx} = 2.97$, $p=0.044$); the bacterial species *G. quercinecans* had larger lesion areas versus the controls, although this was not significant and the $p<0.05$ level ($t_{dunnetx} = 2.79$, $p=0.068$); *B. goodwinii* lesions were smaller and not significantly different from the controls in terms of mean lesion area ($t_{dunnetx} = 1.30$, $p=0.796$), but the necrosis was clearly different to the control wound response (see Figures 1e vs 1f). Inoculation with *Erwinia billingiae*, a known ubiquitous non-pathogenic bacterium, served as a negative control demonstrating lesion area no different to the control wound response. The pattern strengthened when bacteria plus *A. biguttatus* (applied as eggs) were co-inoculated (Figure 1n). Lesions created by *B. goodwinii* and *A. biguttatus* eggs (larvae) were significantly bigger than wound controls ($t_{dunnetx} = 3.40$, $p=0.0125$), as were *G. quercinecans* and *A. biguttatus* ($t_{dunnetx} = 4.65$, $p=0.0002$) and the combination of *B. goodwinii* + *G. quercinecans* + *A. biguttatus* ($t_{dunnetx} = 3.92$, $p=0.0027$) (Figure 1n). Bacteria were re-isolated from lesion margins and at intervals along the larval galleries, which were becoming necrotic forming part of the lesion. Apart from

demonstrating the necrogenic ability of the bacteria, these results showed that the larvae and larval galleries have an important part in increasing lesion area, implicating them in the spread of bacteria within infected trees.

Bacterial positive back-isolation and *Agrilus biguttatus* contamination

There was a high success of back-isolation for both *B. goodwinii* and *G. quercinecans* when these species were included as treatments (70% and 80% respectively); both species were also identified as contaminants (i.e. back-isolated when not part of the treatment inoculation; *B. goodwinii* = 9%; *G. quercinecans* = 16%), although the contamination rate was significantly lower than treatment back-isolation in both cases (*B. goodwinii*: $z = 7.98, p < 0.001$; *G. quercinecans*: $z = 6.65, p < 0.001$). For *G. quercinecans*, the rate of contamination was affected by the presence of *A. biguttatus* eggs: when eggs were absent *G. quercinecans* contamination was 3%; when eggs were present this rose to 38% contamination, a significant increase ($z = 3.59, p < 0.001$). There was no significant effect of *A. biguttatus* eggs on *B. goodwinii* contamination (eggs absent, *B. goodwinii* contamination = 2%; eggs present, *B. goodwinii* contamination = 7%, $z = 1.60, p = 0.11$).

Discussion

Fulfilling Koch's postulates represents the traditional paradigm for proving disease causation. However, contemporary molecular approaches have transformed our appreciation of the role of microbial communities and polymicrobial interactions in disease. There are situations where the one pathogen causes one disease model is not suitable to prove causality and must be adapted to accommodate polymicrobial infections. For example, in decline-diseases of trees, application of Koch's postulates in the strictest sense cannot fully address proving causality of lesion-formation as more than one necrogenic agent is involved in the disease syndrome. Here, causation of AOD lesions was demonstrated using a combined cultivation-based and sequencing approach in a dynamic adaptation of Koch's postulates analogous to that described by Byrd and Segre (2016). Koch's first postulate is fulfilled by the consistent isolation and metagenomic detection of *G. quercinecans*, *B. goodwinii*, and *R. victoriana* in trees affected by AOD.

This combined approach delivered an improved understanding of lesion microbiome composition, as some species were difficult to isolate and culture. For example, *B. goodwinii* and *Lonsdalea quercina* ssp. *britannica* were under-represented, whereas *G. quercinecans*, which is more amenable to isolation and cultivation, was over-represented, with the opposite trend evident in metagenomic studies. Furthermore, *G. quercinecans*, *B. goodwinii* and *R. victoriana* were absent on sites with no history of AOD, were rarely isolated from healthy oak on diseased sites, and had negligible abundance in healthy metagenome samples at AOD sites, complying with Koch's second postulate. Ultra-low levels of *G. quercinecans*, *B. goodwinii* and *R. victoriana* were detected in some of the

512 healthy trees, but only from sites where AOD is present, raising interesting questions
513 about their existence in the wider environment. Possible explanations for the presence of
514 *G. quercinecans*, *B. goodwinii* and *R. victoriana* in healthy trees on AOD sites may
515 include; (1) the tree is in early stages of lesion formation, where visible symptoms have
516 not yet developed, or that (2) *G. quercinecans*, *B. goodwinii* and *R. victoriana* are
517 endophytes or epiphytes that opportunistically multiply after tissue necrosis is initiated by
518 another organism. In the first situation, the chances of detecting asymptomatic lesion
519 formation seem fairly remote, especially as crown condition is not a reliable indicator of
520 tree predisposition status in the early stages of decline development. In the second case,
521 several pieces of evidence counteract the possibility that these organisms multiply
522 opportunistically after tissue necrosis occurs, as (1) we observed the expression of
523 putative necrogenic enzymes and virulence factors of these bacteria in AOD lesions *in*
524 *planta*, implicating them in having an active role in tissue degradation, and (2) *G.*
525 *quercinecans* and *B. goodwinii* caused significant lesion formation in log inoculation
526 trials, demonstrating actual lesion forming capabilities. It is important to note that the oak
527 logs and trees used in our trials were selected from areas with no history of the disease.

528 Furthermore, all three bacterial species possess the genomic capability to cause tissue
529 necrosis, as determined through genomics, *in situ* functional metagenomics and
530 metatranscriptomics. Inoculation onto live oak logs confirmed significant necrogenic
531 ability of *B. goodwinii* and *G. quercinecans*, but further work with *R. victoriana*
532 inoculation is required. Thus, the pathogenic phenotype of *B. goodwinii* and *G.*
533 *quercinecans* has been confirmed, broadly fulfilling Koch's third postulate, that
534 inoculation tests can cause the disease anew. However, bacterial species combinations

caused even greater necrosis indicating cumulative effect and possible synergism, this is a contemporary adaptation of Koch's postulates *sensu stricto*. Finally, once inoculated onto live oak panels, *B. goodwinii* and *G. quercinecans* were re-isolated to fulfil Koch's fourth postulate (although back-isolation of some species was difficult, and there is a need for developing rapid, cost-effective tools to detect pathogens that are difficult to isolate). Consequently, we propose that the biotic component of the AOD lesions is a polybacterial complex comprised primarily of *G. quercinecans* and *B. goodwinii*, which are now established as key causal agents of tissue necrosis, the primary symptom of AOD. Our studies also indicated that other members of the microbiome may contribute to the pathology of AOD. Microbiome analysis suggested that *R. victoriana* is abundant and also important in AOD lesions; however, back-isolation of *R. victoriana* was variable, and further tests to characterize its role and interactions with *B. goodwinii* and *G. quercinecans* are required. In addition, *Lonsdalea quercina* ssp. *britannica* in particular demonstrated variable, but at times virulent pathogenicity in log inoculations, although it was not consistently present in AOD lesions. Our results indicate that *Agrilus* larvae potentiate the spread of these necrogenic bacterial species within tree tissue, generating new break-out points of tissue necrosis, and replicating the observed aetiology of AOD. Ultimately, vascular degradation arising from a combination of bacterial tissue necrosis and inner bark damage from larval galleries exacerbates and accelerates the decline by interrupting carbon resource allocation, preventing accumulation in roots, and reducing water availability.

It is clear that, where possible, microbiome analysis methods together with inoculation assays should become the accepted standard for profiling disease complexes, particularly

when considering complex interactions between microorganisms, insects, the environment, and the host. This phenomenon has never previously been addressed in arboreal systems using the approaches described here. Our work therefore highlights the importance of a systems-level approach for characterizing the pathology of complex diseases, and represents a conceptual and methodological template for adapting Koch's postulates to address the role of microbial communities in disease. In recent years, microbiome studies have transformed our understanding of the role of human gut microbiota in a variety of conditions, including bowel and cardiovascular disease (Frank *et al.*, 2007; Koeth *et al.*, 2013), and consequently, microbiome-wide association studies that link microbial consortia to disease will play an important role in the development of future diagnostics and therapies for disease (Gilbert *et al.*, 2016). For decline-diseases in particular, further adaptation may be required to include the role of host predisposition.

While AOD has been formally described as a major threat to UK oak, similar decline-diseases have been reported in mainland Europe (Biosca *et al.*, 2003; Poza-Carrión *et al.*, 2008; Hartmann *et al.*, 1989; Vansteenkiste *et al.*, 2004; Delatour, 1983), the Middle East, and the Americas (Lynch *et al.*, 2014) indicating that AOD is a global concern that has likely evaded attention due to the complexity of its causative agents. The polybacterial nature of AOD exhibits similarities to other economically important tree diseases such as olive knot disease (Buonaurio *et al.*, 2015) but may also be applied to other complex diseases. Generally, our findings highlight the importance of understanding polymicrobial interactions in the context of future-proofing plant health to protect important but increasingly disease-prone forests and crops that are a fundamental part of our global landscape.

581 **Accession codes**

582 All sequence data described in this study is available under BioProject PRJNA323828
583 and PRJNA321868.

584

585

586

587

588

589

590

591

592

593

594

595

596

597

598

599

References

- Adams AS, Aylward FO, Adams SM, Erbilgin N, Aukema BH, Currie CR, *et al.* (2013). Mountain pine beetles colonizing historical and naive host trees are associated with a bacterial community highly enriched in genes contributing to terpene metabolism. *Appl Environ Microbiol* **79**: 3468–3475.
- Adeolu M, Alnajjar S, Naushad S, Gupta R. (2016). Genome based phylogeny and taxonomy of the ‘*Enterobacteriales*’: proposal for *Enterobacterales* ord. nov. divided into the families *Enterobacteriaceae*, *Erwiniaceae* fam. nov., *Pectobacteriaceae* fam. nov., *Yersiniaceae* fam. nov., *Hafniaceae* fam. nov., *Morganellaceae* fam. nov., and *Budviciaceae* fam. nov. *Int J Syst Evol Microbiol* **66**: 5575–5599.
- Altschul SF, Gish W, Miller W, Myers E, Lipman DJ. (1990). Basic Local Alignment Search Tool. *J Mol Biol* **215**: 403–410.
- Biosca EG, González R, López-López MJ, Soria S, Montón C, Pérez-Laorga E, *et al.* (2003). Isolation and characterization of *Brenneria quercina*, causal agent for bark canker and drippy nut of *Quercus* spp. in Spain. *Phytopathology* **93**: 485–492.
- Boisvert S, Raymond F, Godzaridis E, Laviolette F, Corbeil J. (2012). Ray Meta: scalable *de novo* metagenome assembly and profiling. *Genome Biol* **13**: R122.
- Boyd IL, Freer-Smith PH, Gilligan CA, Godfray HCJ. (2013). The consequence of tree pests and diseases for ecosystem services. *Science* **342**: 823–834.
- Brady C, Denman S, Kirk S, Venter S, Rodríguez-Palenzuela P, Coutinho T. (2010). Description of *Gibbsiella quercinecans* gen. nov., sp. nov., associated with Acute Oak

Decline. *Syst Appl Microbiol* **33**: 444–450.

Brady C, Hunter G, Kirk S, Arnold D, Denman S. (2014). *Rahnella victoriana* sp. nov., *Rahnella bruchi* sp. nov., *Rahnella woolbedingensis* sp. nov., classification of *Rahnella* genomospecies 2 and 3 as *Rahnella variigena* sp. nov. and *Rahnella inusitata* sp. nov., respectively and emended description of the genus *Rahnella*. *Syst Appl Microbiol* **37**: 545–552.

Brady CL, Cleenwerck I, Denman S, Venter SN, Rodríguez-Palenzuela P, Coutinho TA., *et al.* (2012). Proposal to reclassify *Brenneria quercina* (Hildebrand and Schroth 1967) Hauben *et al.* 1999 into a new genus, *Lonsdalea* gen. nov., as *Lonsdalea quercina* comb. nov., descriptions of *Lonsdalea quercina* subsp. *quercina* comb. nov., *Lonsdalea quercina* subsp. *iberica* subsp. nov. and *Lonsdalea quercina* subsp. *britannica* subsp. nov., emendation of the description of the genus *Brenneria*, reclassification of *Dickeya dieffenbachiae* as *Dickeya dadantii* subsp. *dieffenbachiae* comb. nov., and emendation of the description of *Dickeya dadantii*. *Int J Syst Evol Microbiol* **62**: 1592–1602.

Brown L, Li X. (2005). Confidence intervals for two sample binomial distribution. *J Stat Plan Inference* **130**: 359–375.

Brown N, Inward DJG, Jeger M, Denman S. (2015). A review of *Agrilus biguttatus* in UK forests and its relationship with acute oak decline. *Forestry* **88**: 53–63.

Brown N, Jeger M, Kirk S, Xu X, Denman S. (2016). Spatial and temporal patterns in symptom expression within eight woodlands affected by Acute Oak Decline. *For Ecol Manage* **360**: 97–109.

Brown N, Jeger MJ, Kirk S, Williams DT, Xu X, Pautasso M, *et al.* (2017). Acute Oak

643 Decline and *Agrilus biguttatus*: the co-occurrence of stem bleeding and D-shaped
644 emergence holes in Great Britain. *Forests* **8**: 87.

645 Buonauro R, Moretti C, da Silva DP, Cortese C, Ramos C, Venturi V. (2015). The olive
646 knot disease as a model to study the role of interspecies bacterial communities in plant
647 disease. *Front Plant Sci* **6**: 434.

648 Byrd AL, Segre JA. (2016). Adapting Koch's postulates. *Science* **351**: 224–226.

649 Charkowski A, Blanco C, Condemine G, Expert D, Franza T, Hayes C, *et al.* (2012). The
650 role of secretion systems and small molecules in soft-rot *Enterobacteriaceae*
651 pathogenicity. *Annu Rev Phytopathol* **50**: 425–449.

652 Clarke K, Gorley R. (2015). PRIMER v7: User Manual/Tutorial. PRIMER-E. Plymouth.

653 Delatour C. (1983). Die-back of oak in Europe. *Biol For* **35**: 265–282.

654 Cohen W, Yang Z, Stehman SV, Shroeder TA, Bell DM, Masek JG, *et al.* (2016). Forest
655 disturbance across the conterminous United States from 1985-2012: The emerging
656 dominance of forest decline. *For Ecol Manage* **360**: 242-252.

657 Denman S, Brady C, Kirk S, Cleenwerck I, Venter S, Coutinho T, *et al.* (2012).
658 *Brenneria goodwinii* sp. nov., associated with acute oak decline in the UK. *Int J Syst Evol*
659 *Microbiol* **62**: 2451–2456. Denman S, Brown N, Kirk S, Jeger M, Webber J. (2014). A
660 description of the symptoms of Acute Oak Decline in Britain and a comparative review
661 on causes of similar disorders on oak in Europe. *Forestry* **87**: 535–551.

662 Denman S, Plummer S, Kirk S, Peace A, McDonald JE. (2016). Isolation studies reveal a
663 shift in the cultivable microbiome of oak affected with Acute Oak Decline. *Syst Appl*

664 *Microbiol* **39**: 18–20.

665 Denman S, Webber J. (2009). Oak declines: new definitions and new episodes in Britain.

666 *Q J For* **103**: 285–290.

667 Falkow S. (1988). Molecular Koch’s postulates applied to microbial pathogenicity. *Rev*

668 *Infect Dis* **10**: 7–10.

669 Frank DN, St Amand AL, Feldman RA, Boedeker EC, Harpaz N, Pace NR. (2007).

670 Molecular-phylogenetic characterization of microbial community imbalances in human

671 inflammatory bowel diseases. *Proc Natl Acad Sci U S A* **104**: 13780–13785.

672 Fredricks DN, Relman DA. (1996). Sequence-based identification of microbial

673 pathogens: a reconsideration of Koch’s postulates. *Clin Microbiol Rev* **9**: 18–33.

674 Gibbs JN, Greig BJW. (1997). Biotic and abiotic factors affecting the dying back of

675 pedunculate oak *Quercus robur* L. *Forestry* **70**: 399–406.

676 Gilbert JA, Quinn RA, Debelius J, Xu ZZ, Morton J, Garg N, *et al.* (2016). Microbiome-

677 wide association studies link dynamic microbial consortia to disease. *Nature* **535**: 94–

678 103.

679 Gower JC. (1966). Some distance properties of latent root and vector methods used in

680 multivariate analysis. *Biometrika* **53**: 325–338.

681 Hartmann G, Blank R, Lewark S. (1989). Oak Decline in Northern Germany:

682 distribution, symptoms, possible causes. *Forst und holz* **44**: 475–487.

683 Hildebrand D, Schroth M. (1967). A new species of *Erwinia* causing the drippy nut

684 disease of live oaks. *Phytopathology* **57**: 250–253.

685 Hill AB. (1965). The environment and disease: Association or causation? *Proc R Soc*
686 *Med* **58**: 295–300.

687 Joshi N, Fass J. (2011). Sickle. A sliding-window, adaptive, quality-based trimming tool
688 for FastQ files. [https:// github.com/najoshi/sickle](https://github.com/najoshi/sickle)

689 Koeth RA, Wang Z, Levison BS, Buffa JA, Org E, Sheehy BT, *et al.* (2013). Intestinal
690 microbiota metabolism of L-carnitine, a nutrient in red meat, promotes atherosclerosis.
691 *Nat Med* **19**: 576–585.

692 Lamichhane JR, Venturi V. (2015). Synergisms between microbial pathogens in plant
693 disease complexes: a growing trend. *Front Plant Sci* **6**: 385.

694 Lynch SC, Zambino PJ, Scott TA, Eskalen A. (2014). Occurrence, incidence and
695 associations among fungal pathogens and *Agrilus auroguttatus*, and their roles in
696 *Quercus agrifolia* decline in California. *For Pathol* **44**: 62–74.

697 Manion P. (1981). Tree Disease Concepts. Prentice Hall, Englewood Cliffs: New Jersey.

698 Martin M. (2011). Cutadapt removes adapter sequences from high-throughput sequencing
699 reads. *EMBnet.journal* **17**: 10.

700 McDowell NG, Beerling DJ, Breshears DD, Fisher RA, Raffa KF, Stitt M. (2011). The
701 interdependence of mechanisms underlying climate-driven vegetation mortality. *Trends*
702 *Ecol Evol* **26**: 523–532.

703 Meyer F, Paarmann D, D’Souza M, Olson R, Glass E, Kubal M, *et al.* (2008). The
704 metagenomics RAST server – a public resource for the automatic phylogenetic and
705 functional analysis of metagenomes. *BMC Bioinformatics* **9**: 386.

706 Millar CI, Stephenson NL. (2015). Temperate forest health in an era of emerging
707 megadisturbance. *Science* **349**: 823–826.

708 Minot SS, Krumm N, Greenfield NB. (2015). One Codex : A sensitive and accurate data
709 platform for genomic microbial identification. *bioRxiv*. e-pub ahead of print, doi:
710 10.1101/027607.

711 Moran F, Seiichi N, Starr M. (1968). Oligogalacturonide trans-Eliminase of *Erwinia*
712 *carotovora*. *Arch Biochem Biophys* **125**: 734–741.

713 Oliva J, Stenlid J, Martínez-Vilalta J. (2014). The effect of fungal pathogens on the water
714 and carbon economy of trees: Implications for drought-induced mortality. *New Phytol*
715 **203**: 1028–1035.

716 Parks DH, Tyson GW, Hugenholtz P, Beiko RG. (2014). STAMP: Statistical analysis of
717 taxonomic and functional profiles. *Bioinformatics* **30**: 3123–3124.

718 Pautasso M, Schlegel M, Holdenrieder O. (2015). Forest health in a changing world.
719 *Microb Ecol* **69**: 826–842.

720 Poza-Carrión C, Aguilar I, Gallego FJ, Nuñez-Moreno Y, Biosca EG, González R, *et al.*
721 (2008). *Brenneria quercina* and *Serratia* spp. isolated from Spanish oak trees: Molecular
722 characterization and development of PCR primers. *Plant Pathol* **57**: 308–319.

723 Rackham O. (2008). Ancient woodlands: Modern threats. *New Phytol* **180**: 571–586.

724 Reed K. (2016). The role of the two spotted oak buprestid in Acute Oak Decline. *Pract*
725 *Bull Chart Inst Ecol Environ Manag* **91**: 33–36.

726 Sallé A, Nageleisen LM, Lieutier F. (2014). Bark and wood boring insects involved in

727 oak declines in Europe: Current knowledge and future prospects in a context of climate
 728 change. *For Ecol Manage* **328**: 79–93.

729 Sapp M, Lewis E, Moss S, Barrett B, Kirk S, Elphinstone J, *et al.* (2016). Metabarcoding
 730 of bacteria associated with the Acute Oak Decline syndrome in England. *Forests* **7**: 95.

731 Seemann T. (2014). Prokka: rapid prokaryotic genome annotation. *Bioinformatics* **30**:
 732 2068–2069.

733 Sinclair WA. (1965). Comparisons of recent declines of white ash, oaks, and sugar maple
 734 in northeastern woodlands. *Cornell Plant* **20**: 62–67.

735 Sinclair W, Lyon H. (2005). Diseases of trees and shrubs. Comstock Publishing
 736 Associates, Ithaca: New York.

737 Tampakaki AP, Skandalis N, Gazi AD, Bastaki MN, Sarris PF, Charova SN, *et al.*
 738 (2010). Playing the Harp: Evolution of our understanding of *hrp/hrc* genes. *Annu Rev*
 739 *Phytopathol* **48**: 347–370.

740 Thomas F. (2008). Recent advances in cause-effect research on oak decline in Europe.
 741 *CAB Rev Perspect Agric Vet Sci Nutr Nat Resour* **3**: 1–12.

742 Toth IK, Pritchard L, Birch PRJ. (2006). Comparative genomics reveals what makes an
 743 enterobacterial plant pathogen. *Annu Rev Phytopathol* **44**: 305–336.

744 Vansteenkiste D, Tirry L, Van Acker J, Stevens M. (2004). Predispositions and
 745 symptoms of *Agilus* borer attack in declining oak trees. *Annu For Sci* **61**: 815–823.

746 Versluis D, Leimena MM, Garcia JR, Andrea MMD. (2015). Mining microbial
 747 metatranscriptomes for expression of antibiotic resistance genes under natural conditions.

748 *Sci Rep* **5**: 11981.

749 Vinatzer BA, Monteil CL, Clarke CR. (2014). Harnessing population genomics to
750 understand how bacterial pathogens emerge, adapt to crop hosts, and disseminate. *Annu*
751 *Rev Phytopathol* **52**: 19–43.

752 Wood DE, Salzberg SL. (2014). Kraken: ultrafast metagenomic sequence classification
753 using exact alignments. *Genome Biol* **15**: R46.

754 Yao J, Chen QL, Shen AX, Cao W, Liu YH. (2013). A novel feruloyl esterase from a soil
755 metagenomic library with tannase activity. *J Mol Catal B Enzym* **95**: 55–61.

756

Acknowledgments: This work was supported by Woodland Heritage, the Rufford Foundation, the Monument Trust, the JPG fund, HPC Wales, the Forestry Commission and DEFRA (TH0108). We thank the private land owners, public forest estate, the National Trust and the Woodland Trust for site access and allowing tissue sampling of oaks. We are also grateful to the internal reviewers for their helpful suggestions on the manuscript. Sequence data generated in this study is available under NCBI BioProject PRJNA323828. We are grateful to Katy Reed and Daegan Inward for supplying *Agrilus* eggs, the Forest Research Technical Service Unit for supplying logs and assistance with field and log trials. We are grateful to Lucille Rainbow and Margaret Hughes of the Centre of Genomic Research for library preparation and sequencing.

Supplementary Information accompanies this paper on The ISME Journal website

Fig 1. Symptoms of Acute Oak Decline observed in the field (a-d), reproducing the symptoms of AOD through log inoculations with combinations of bacteria and *Agrilus* larvae associated with AOD (e-l), and statistical analysis of lesion formation in log inoculations (m-n). (a) Stem bleeds symptomatic of Acute Oak Decline (AOD) on a mature *Quercus robur* in the field. (b) Close up view of stem bleeds and bark cracks characteristic of AOD. (c) Cross-section through AOD stem bleeds, showing degradation of vascular tissue (red arrow). (d) Longitudinal section through an AOD stem bleed showing the association between bacterial lesions (blue arrow) and *Agrilus biguttatus* galleries (red arrow). (e) Wound response in the innerbark of the control treatment (inoculation of a wound with sterile water) of log inoculation trials. (f) Lesion formed in the innerbark of oak logs inoculated with *Brenneria goodwinii* in log inoculation trials. (g) Lesion formed in the innerbark of oak logs inoculated with *Gibbsiella quercinecans* in log inoculation trials. (h) Lesion formed in the innerbark of oak logs inoculated with a combination of *B. goodwinii* and *G. quercinecans* in log inoculation trials. (i) Wound response in the innerbark of the *A. biguttatus* treatment (inoculation of a wound with live eggs of *A. biguttatus*). Note the wound response (blue arrow) and clean gallery created by a larva (red arrow). (j) Lesion formed in the innerbark of oak logs inoculated with *B. goodwinii* and eggs of *A. biguttatus*. Note the lesion developing from the inoculation point (blue arrow), and from the galleries (red arrow). (k) Lesion formed in the innerbark of oak logs inoculated with *G. quercinecans* plus eggs of *A. biguttatus*. Note the lesion developing from the inoculation point (blue arrow), and from the galleries (red arrow). (l) Lesion formed in the innerbark of oak logs inoculated with *B. goodwinii*, *G. quercinecans* plus eggs of *A. biguttatus*. Note the lesion developing from the inoculation point (blue arrow), and from the galleries (red arrow). (m) Mean lesion area formed in the inner bark of logs inoculated with bacteria. Colour indicates statistical significance (see key). The bacterial species inoculated were back-isolated in fulfilment of Koch's fourth postulate. Error bars = - standard error. (n) Mean lesion area formed in the inner bark of logs inoculated with bacteria and *A. biguttatus* eggs. Colour indicates statistical significance (see key in m). The bacterial species inoculated were back-isolated in fulfilment of Koch's fourth postulate. *Eb* – *Erwinia billingiae*, *Bg* – *Brenneria goodwinii*, *Gq* – *Gibbsiella quercinecans*. Error bars = - standard error.

Fig 2. Comparative metagenomic analysis of the taxonomic composition and function of the oak microbiome in healthy tissue (from trees without AOD) and diseased tissue (from trees with AOD). (a) The plot depicts the genera represented in all metagenomes based on One Codex binning of raw reads, demonstrating a clear shift in the microbial community between healthy and diseased trees. (b) *Brenneria* and *Gibbsiella* are statistically correlated to diseased tissue, as shown by a principal coordinate ordination analysis based on statistics calculated by Primer v7 and PERMANOVA+ using One Codex binning data. The first two axes depict the plotted community composition. Correlation vectors in the graph are significant using an $R^2 > 0.05$. A Welch's *t*-test was performed to test significance of differences between key taxa (identified above) between healthy and diseased trees (pooled abundances for each factor). Resultant *p*-values from Welch's *t*-test are overlaid on the correlation biplot. (c) Gene groups involved in bacterial phytopathogenic activity are significantly increased in AOD diseased trees, as shown by comparative functional analysis of SEED subsystem categories as annotated by MG-RAST on assembled metagenome contigs. The analysis of SEED subsystem metagenome data was performed using Stamp. Statistically significant functional differences between diseased and healthy communities were calculated using a G-test with Yates' correction. The Newcombe-Wilson test was performed to calculate confidence intervals between two binomial population proportions. (d) The genomes of 17 species were found to be common across all AOD metagenomes. Visualization of Kraken metagenome analysis of stem samples demonstrates the shifts in bacterial microbiome compositions. Bubble sizes are categorized based on the relative percentage frequencies of Kraken species-level alignment of raw metagenome sequencing reads, and are depicted in the figure as 1 (read frequency <0.01), 2 (read frequency 0.01-0.1), 3 (read frequency 0.1-1), 4 (read frequency 1-10) and 5 (read frequency >10). Red bubbles signify samples from diseased trees, while blue bubbles signify samples from healthy trees. The 17 common species were determined based on common occurrence across all diseased tissue samples. Additionally, six species more abundant among the healthy trees were included to provide a contrasting shift.

Fig 3. Functional genome analysis of *Gibbsiella quercinecans* FRB97 (T), *Brenneria goodwinii* FRB141 (T), and *Rahnella victoriana* BRK18a. (a) Circular representations of *B. goodwinii* FRB141, *G. quercinecans* FRB97, and *R. victoriana* BRK18a genomes. From outside to inside, circles represent: (1) Assembled bacterial genomes, outermost (orange) circle, with encoded secretion system annotated at their genomic loci. (2) Metatranscriptome heatmap. Alignment of two *in silico* combined metatranscriptomes recovered from two necrotic lesions of AOD-affected trees, against the bacterial genomes. Blue saturation represents increasing transcript alignments. (3-9) Seven metagenomes from necrotic lesions on AOD affected trees and one healthy tree (metagenomes were extracted from two sites, Attingham and Runs Wood), were aligned through their coding domains to homologous regions in the bacterial genomes. (3) Attingham healthy (AT1) aligned metagenome coding domains (light purple). (4) Attingham diseased (AT7) aligned metagenome coding domains (aqua). (5) Attingham diseased (AT8) aligned metagenome coding domains (blue). (6) Attingham diseased (AT9) aligned metagenome coding domains (orange). (7) Runs Wood diseased (RW1) aligned metagenome coding domains (green). (8) Runs Wood diseased (RW2) aligned metagenome coding domains (pink). (9) Runs Wood diseased (RW3) aligned metagenome coding domains (grey). (10) G+C content across the bacterial genome. (11) G+C skew across the bacterial genome. (b) Schematic diagram of transporters and transported proteins recovered from lesion metatranscriptomes and aligned against each bacterial genome. Number of significantly expressed (>3 aligned transcripts, covering >20% of the gene) genes is shown in parentheses. (c) Number of transcripts aligned against selected virulence genes encoded within *B. goodwinii* FRB141, *G. quercinecans* FRB97, and *R. victoriana* BRK18a. Gene categories are represented by the following colours, red - plant cell wall degrading enzymes (PCWDE), purple - general secretory pathway (GSP), yellow - type II secretion system (T2SS), blue - type III secretion system (T3SS), pink - type III secretion system effectors (T3SS effectors), and green - global regulators (GR)

853 **Supplemental Tables:**

854 **Supplementary Table S1. Forest and woodland sites used for bacterial isolation from**
855 **diseased (Acute Oak Decline) and healthy oak trees.** Ordinance survey co-ordinates are
856 provided.

857 **Supplementary Table S2. Annual log inoculation test treatments using single and**
858 **multiple bacterial species inoculations, with or without the addition of eggs of *Agrilus***
859 ***biguttatus*.** Microbiome samples for metagenomic analysis were gathered at three of these
860 sites, two sites in Attingham park were sampled on two dates: Samples AT1, AT7, AT8 and
861 AT9 were collected from OS Eastings 356033, Northings 310372, in November 2013 as part
862 of the CSR framework of sample collection; and samples AT2, AT3, AT4, AT5 and AT6 at
863 OS Eastings 356033, Northings 310372, were collected additionally in June 2015. Samples
864 RW1, RW2 and RW3 were collected from Runs Wood in February 2014 (OS Eastings
865 563207, Northings 310858) as part of the CSR framework, and samples ROW1, ROW2 and
866 ROW3 were additional samples collected from Ross-on-Wye (OS Eastings 357887,
867 Northings 221731), in June 2015. Ordinance survey co-ordinates are provided.

868 **Supplementary Table S3. Accession numbers for genomic, metagenomic,**
869 **metatranscriptomic libraries.** All libraries were submitted to the National Center for
870 Biotechnology Information. *Genome assembly.

871 **Supplementary Table S4. Alignment of metagenome coding domains to the genomes of**
872 ***Brenneria goodwinii* FRB141 (T), *Gibbsiella quercinecans* FRB97 (T), and *Rahnella***
873 ***victoriana* BRK18a (T) and two control genomes, *Pectobacterium carotovorum* ssp.**
874 ***carotovorum* PC1, a known plant pathogen, and the endophyte, *Paenibacillus polymyxa***
875 **SC2.** To assess the abundance of *Brenneria goodwinii*, *Gibbsiella quercinecans*, and
876 *Rahnella victoriana* in the Acute Oak Decline (AOD) lesion metagenome, alignments were
877 made between assembled metagenomic libraries from AOD diseased and a healthy oak tree
878 (AT1), and assembled bacterial coding domains of *B. goodwinii*, *G. quercinecans* and *R.*
879 *victoriana*. In addition, to assess the veracity of our alignment approach, we tested two
880 additional control genomes; *Pectobacterium carotovorum* ssp. *carotovorum* PC1 is a well-
881 characterized bacterial phytopathogen that has not previously been associated with oak. This
882 genome would therefore not be expected to be present within AOD lesion metagenomes, and
883 therefore provides an indication of the potential level of alignment matches from homologous
884 genes found within other members of the *Enterobacteriaceae*. Only a small proportion of

metagenome coding domains (44-183) mapped with *P. carotovorum* ssp. *carotovorum* PC1, suggesting that the greater proportion of matches that occur between *B. goodwinii*, *G. quercinecans* and *R. victoriana* reflect the actual presence and abundance of those species in AOD lesion metagenomes, rather than them representing metagenome coding domains from other *Enterobacteriaceae* that have been falsely attributed to our focal species (*B. goodwinii*, *G. quercinecans* and *R. victoriana*). *Paenibacillus polymyxa* SC2 is a putative endophyte found at low relative abundance within symptomatic and healthy oak tissue and was included as a control to ensure that the high numbers of metagenome coding domains mapped to *B. goodwinii*, *G. quercinecans* and *R. victoriana* are accurate and would not occur in the same way for other *Enterobacteriaceae* that are known to also be present in AOD lesions. The low numbers of coding domains that map against *P. polymyxa* SC2 (0-8) again supports the veracity of our alignment approach, and the abundance of *B. goodwinii*, *G. quercinecans* and *R. victoriana* in AOD lesion tissue.

Supplementary Table S5. Combined Kraken output data table. Output data of Kraken metagenomic binning of all metagenomes. Where U=unidentified, D=domain, P=phylum, C=class, O=order, F=family, G=genus, S=species. The standard Kraken database was supplemented using the sequenced genomes of *Rahnella victoriana*, *Lonsdalea quercina*, *Gibbsiella quercinecans* and *Brenneria goodwinii*.

Supplementary Table S6. Occurrence of the most commonly isolated bacterial taxa on sites and trees.

Supplementary Table S7. Significance of genera detected in healthy and symptomatic metagenome samples. Welch's unequal variances *t*-tests of genus abundance detected in metagenome datasets derived from healthy and diseased oak trees. ^aAsymptotically F distributed.

Supplementary Table S8. Genome metrics of *Brenneria goodwinii* FRB141 (T), *Gibbsiella quercinecans* FRB97 (T), and *Rahnella victoriana* BRK18a (T) sequenced bacterial isolates. Bacteria were isolated from necrotic lesions of Acute Oak Decline affected trees.

Supplementary Table S9. Genomic loci of putative bacterial virulence genes. Genes encoded within *Brenneria goodwinii* FRB141, *Gibbsiella quercinecans* FRB97, and *Rahnella victoriana* BRK18a. PCWDE – plant cell wall degrading enzymes, T2SS – type II secretion system, T3SS – type III secretion system, T4SS – type IV secretion system.