

Host-microbiota-insect interactions drive emergent virulence in a complex tree disease

Doonan, James; Broberg, Martin; Denman, Sandra; McDonald, James

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- 1 Host-microbiota-insect interactions drive emergent virulence in a
- 2 complex tree disease
- 3
- 4 James M. Doonan^{1,2}, Martin Broberg^{1,3}, Sandra Denman⁴ and James E. McDonald^{1*}
- ⁵ ¹School of Natural Sciences, Bangor University, Deiniol Road, Bangor, Gwynedd, LL57
- 6 2UW, UK.
- ⁷ ²Department of Geosciences and Natural Resource Management, University of Copenhagen,
- 8 Rolighedsvej 23, 1958 Frederiksberg C, Denmark.
- ⁹ ³Faculty of Biological and Environmental Sciences, University of Helsinki, Finland.
- ⁴Forest Research, Centre for Forestry and Climate Change, Alice Holt Lodge, Farnham,
- 11 Surrey GU10 4LH, UK.
- 12 *Corresponding author

13 Abstract

14 Forest declines caused by climate disturbance, insect pests and microbial pathogens threaten 15 the global landscape, and tree diseases are increasingly attributed to the emergent properties 16 of complex ecological interactions between the host, microbiota and insects. To address this 17 hypothesis, we combined reductionist approaches (single and polyspecies bacterial cultures) 18 with emergentist approaches (bacterial incoluations in an oak infection model with the 19 addition of insect larvae) to unravel the gene expression landscape and symptom severity of 20 host-microbiota-insect interactions in the Acute Oak Decline (AOD) pathosystem. AOD is a 21 complex decline disease characterised by predisposing abiotic factors, inner bark lesions 22 driven by a bacterial pathobiome, and larval galleries of the bark-boring beetle Agrilus 23 biguttatus. We identified expression of key pathogenicity genes in Brenneria goodwinii, the dominant member of the AOD pathobiome, tissue-specific gene expression profiles, 24 25 cooperation with other bacterial pathobiome members in sugar catabolism, and demonstrated 26 amplification of pathogenic gene expression in the presence of Agrilus larvae. This study 27 highlights the emergent properties of complex host-pathobiota-insect interactions that 28 underlie the pathology of diseases that threaten global forest biomes.

29 Introduction

30 Global forests provide essential ecological, economic and cultural services, but their capacity 31 for carbon storage and climate regulation is increasingly threatened by altered climatic 32 conditions and increased attack by pests and pathogens [1,2]. In recent decades, devastating 33 outbreaks of tree disease such as chestnut blight [3], Dutch elm disease [4], and ash dieback 34 [5], have changed the global landscape, and tree pests and diseases therefore represent a 35 major future threat to forest biomes. Such diseases often involve the activity of both insect 36 pests and microbial pathogens, and ultimately arise from complex interactions between the 37 host, environment, pests and pathogens [6-8].

38 Acute Oak Decline (AOD) is a complex decline disease mediated by abiotic predisposing 39 factors (temperature, rainfall, nutrients) [9] and biotic contributing factors (insect and 40 bacterial) [8] that are a major threat to native oak in the UK, with similar declines described 41 in continental Europe [10–12], Asia [13] and America [14]. The characteristic disease 42 symptoms are outer bark cracks with dark exudates (bleeds), which overlie necrotic tissue in 43 the inner bark, and larval galleries and exit holes of the two-spotted buprestid beetle Agrilus 44 biguttatus [10]. Previously, we demonstrated that tissue necrosis on AOD affected trees is 45 caused by a polybacterial complex (pathobiome) which macerates pectin connective tissue 46 within the cells, resulting in inner bark lesions on oak stems [8,15]. The pathobiome is a 47 complex assemblage of organisms that combine to cause disease in host organisms and 48 challenge strict adherence to Koch's postulates [16]. It has previously been shown that AOD 49 is not caused by a single pathogen, but results from interactions between the pathobiome, A. 50 biguttatus, the host and it's environment [8]. Within the AOD pathobiome several bacteria 51 are consistently identified, primarily Brenneria goodwinii, Gibbsiella quercinecans, Rahnella 52 victoriana, and occasionally, Lonsdalea britannica.

Brenneria goodwinii, is the most active member of the lesion pathobiome, and is thought to be the primary agent of bacterial canker in AOD [8,15]. *Agrilus* larvae are also associated with AOD lesions, and spread necrogenic members of the pathobiome through the inner bark tissue, amplifying the area of tissue necrosis in the inner bark [8].

57 Unravelling the mechanistic processes and complex multidimensional interactions between 58 the host, environment, insects, and the pathobiome that underlie the aetiology of complex tree 59 diseases is challenging, but represents a major knowledge gap. Considering pathobiome 60 virulence as an emergent property [17], where emerging properties cannot be explained by 61 their individual components and are greater than the sum of their individual components, is therefore an attractive framework in conceptualising complex tree diseases. Here, we 62 63 hypothesise that host-microbiota-insect interactions combine to cause emergent properties of 64 pathobiome virulence in AOD. To investigate this, we combined reductionist approaches 65 (interactions with oak tissue in single and polyspecies bacterial culture) with emergentist 66 approaches (bacterial inoculations in an oak infection model with the addition of insect 67 larvae) to unravel the gene expression landscape of host-microbiota-insect interactions in the 68 Acute Oak Decline (AOD) pathosystem.

69

70 Results and Discussion

Inoculation of *B. goodwinii*, *G. quercinecans* and *L. britannica* onto oak logs with *A. biguttatus* eggs

73 Oak logs were inoculated with either B. goodwinii, G. quercinecans or L. britannica (single, 74 bacteria-only treatments), or in combination with A. biguttatus eggs (single bacterial species 75 plus Agrilus treatments) (i.e. six treatments). RNAseq analysis of the resultant stem lesions 76 (or 'clean' stem tissue, for control treatments) revealed that apart from host genes, B. 77 goodwinii genes were the most actively expressed amongst the bacterial species tested (figure 78 1 and electronic supplementary material table, S1), concurring with previous results [8,15]. 79 The complete genome of *B. goodwinii* FRB141 contains 4869 genes and the highest levels of 80 B. goodwinii gene activity in the log infection tests were detected in treatments where B. 81 goodwinii was co-inoculated with A. biguttatus eggs (515, 3924, and 2464 genes expressed in 82 each replicate, respectively) and there was positive detection of *B. goodwinii* via RT-qPCR 83 (please see materials and methods for our definition of active genes briefly, these are genes 84 which were not differentially expressed, but were deemed 'active' as they passed expression 85 filters e.g. Transcripts per Million, but differ from subsequent analyses that focussed on 86 differential gene expression). By comparison, only one of the three *B. goodwinii* only 87 inoculations produced necrosis, with 3819 active genes detected, while the other two 88 inoculations did not show appreciable lesion development and only 88 and 96 active genes 89 were detected.

Lesions barely developed in *L. britannica* inoculations, with low activity (11 +/- 3 active genes) detected, although this species was previously isolated from naturally symptomatic material and has the genomic potential to cause tissue necrosis [18]. By comparison, when co-inoculated with *A. biguttatus* eggs, two of the three inoculations developed dramatic, typical AOD lesions, with 46 and 1607 *L. britannica* genes active (figure 1), but both *B. goodwinii* and *G. quercinecans* were also reisolated via RT-qPCR, and 852 and 2942 *B. goodwinii* genes and 579 and 320 *G. quercinecans* genes were found to be active. Notably, *G. quercinecans* which has been consistently isolated from environmental AOD lesions and can cause necrotic lesions on oak [8], had low activity in log inoculations (143 +/- 71 active genes), but had higher gene activity and significant lesion formation when combined with *A. biguttatus* (444 +/- 225).

101 Thus, with the exception of a single B. goodwinii inoculation, none of the single isolate 102 inoculations created significant lesions or demonstrated high gene expression, which supports 103 our hypothesis that although these organisms can be pathogenic, emergent virulence is 104 dependent upon complex host-pathobiome-insect interactions. However, when co-inoculated 105 with A. biguttatus eggs that developed into larvae, typical AOD symptoms were developed 106 and B. goodwinii gene activity was highly increased. This suggests that the presence of A. 107 biguttatus larvae provides a stimulus for enhanced B. goodwinii pathogenicity. Furthermore, 108 the biggest lesions formed when genes of all three bacterial species were detected. Despite the fact that only single species inoculations were made, the occurrence of B. goodwinii and 109 110 G. quercinecans in the L. britannica plus Agrilus treatment could be explained either by the 111 bacteria already being present as endosymbionts of the non-symptomatic oak logs, or by 112 them gaining entry through wound inoculations, or that A. biguttatus is a vector of B. 113 goodwinii, either incidentally or that it resides within A. biguttatus as part of the microbiome 114 and is deposited when feeding or egg laying [10]. This suggests that the presence of A. 115 biguttatus larvae provides a stimulus for enhanced B. goodwinii pathogenicity. However, 116 there is no previous evidence showing that A. biguttatus is a vector of B. goodwinii, G. 117 quercinecans or L. britannica and the bacteria-beetle relationship may be as co-infecting 118 agents taking advantage of declining oak trees [19].

119 Our results demonstrate that the driver of variation between non-symptomatic and 120 symptomatic oak trees was bacterial inoculum (P = 0.031) and the prescence of A. biguttatus 121 larvae (P = 0.005) (figure 1). Possible sources of variation in gene activity between 122 symptomatic and non-symptomatic trees were tested in a multivariate model, these were: 123 actual lesion size, presence or absence of A. biguttatus, bacterial inoculum, and between 124 replicate differences. Biological replicates and lesion size did not account for significant 125 variation in gene activity (P > 0.05). Furthermore, differential gene expression analysis 126 revealed that the number of genes expressed in G. quercinecans and L. britannica was

relatively small, whereas in *B. goodwinii* inoculations, a substantial portion of the *B. goodwinii* geneset (electronic supplementary material table, S2) was differentially expressed.
Therefore, the following differential gene expression analysis of *B. goodwinii* was directly
compared against control treatments and *B. goodwinii* when co-inoculated with *A. biguttatus*larvae.

132

B. goodwinii has a high number of significantly upregulated genes in log inoculations when inoculated with A. biguttatus

135 Differential gene expression analysis of *B. goodwinii* log inoculations (bacteria only) 136 compared against wound and water controls revealed 191 genes were significantly 137 differentially upregulated (electronic supplementary material table, S2). Comparison of the B. 138 goodwinii and A. biguttatus treatment with the wound control resulted in 552 upregulated B. 139 goodwinii genes. Variance between expressed genes within transcriptomic datasets was 140 measured using principal component analyses (PCA) (figure 2a). This PCA collapsed 73% of 141 the variance and revealed clear separation between transcipt abundance in B. goodwinii 142 infected oak logs compared to the control (figure 2a, bottom). The same pattern was found in 143 B. goodwinii and A. biguttatus inoculated oak logs where 81% of the variance was captured 144 in a PCA and revealed distinct expression patterns in comparison to oak control logs (figure 145 2a, top). Analysis of differential expression of gene families, revealed significant upregulation of putative pathogenic families in B. goodwinii and A. biguttatus egg inoculations when 146 147 compared to *B. goodwinii* only oak logs. These gene families were identified using geneset 148 enrichment analysis and revealed that gene families were upregulated in *B. goodwinii* by the 149 presence of A. biguttatus eggs. Significantly upregulated functional groups include bacterial 150 pathogenicity homologs, such as bacterial secretion systems (P=0.04, KEGG family 03070), 151 terpenoid biosynthesis (P=0.04, KEGG family 00130), biofilm formation (P=0.007, KEGG 152 family 02026), and quorum sensing (P=0.01, KEGG family 02024) (figure 2b). Differential 153 gene expression analysis between oak log incoulations revealed significant upregulation of 154 pathogenicity associated genes in B. goodwinii and A. biguttatus oak logs compared to 155 control, in comparison to differential expression of the same gene in *B. goodwinii* only oak 156 logs when compared to contol. Genes were functionally annotated using homologs in closely 157 related bacteria (see methods). Significantly upregulated functional homologs included a biofilm formation gene, exoglucanase B - chvB (Padj < 0.0001 in B. goodwinii + A. 158 159 biguttatus vs. healthy, compared to B. goodwinii only vs. healthy, which had no P value due 160 to low transcript expression), an adherence gene - fhaB (Padj=0.03 in B. goodwinii + A. 161 *biguttatus* vs. healthy, compared to P = 0.02, N.B *Padj* was NA as the mean read count was 162 low in *B. goodwinii* only v healthy), $poly(\beta$ -D-mannuronate) C5 epimerase 1, a biofilm 163 formation and quorum sensing gene - algG (Padj < 0.0001 in B. goodwinii + A. biguttatus v 164 healthy, compared to Padj = 0.0006 B. goodwinii only vs. healthy). Poly(β -D-mannuronate) C5 epimerase 1 is a large, type I secreted adhesin which is found in shiga toxin producing E. 165 166 coli strains and in disease formation of the bacterial phytopathogen Pectobacterium 167 *atrosepticum* [20,21]. Both exoglucanase B and poly(β-D-mannuronate) C5 epimerase 1 were 168 significantly upregulated in B. goodwinii and G. quercinecans only live log inoculations 169 indicating that A. biguttatus may not be the only stimulus for its expression. The actual 170 stimulus may be carried by A. biguttatus or may reside in the wider environment. Similar to 171 the type I secreted proteins, two copies of the two-partner secreted filamentous hemagglutinin 172 (*fhaB*), a bacterial virulence gene were expressed by *B. goodwinii* across live log 173 transcriptomes. As described above, the number of genes expressed in *B. goodwinii* when *A*. 174 biguttatus was present was greater than B. goodwinii only inoculations (191 vs. 552, 175 respectively), but in addition the number of pathogenic gene homologs expressed increased 176 when A. biguttatus eggs were combined with B. goodwinii (figure 2c).

177 The T3SS is a primary virulence factor in seven of the top ten bacterial plant pathogens [22]. 178 B. goodwinii encodes a complete T3SS and multiple effectors, which is likely to be a key 179 pathogenicity component within AOD tissue necrosis [18]. Within B. goodwinii and A. 180 biguttatus live log inoculations, four T3 effectors are significantly differentially expressed, 181 only one of which is expressed in *B. goodwinii* only inoculations (figure 2c). Significantly 182 expressed T3 effectors are; HopPtoL (Padj = 0.02), SrfB (Padj = 0.02), AvrE_2 (Padj = 183 0.015), in addition to AvrE_1 which is significantly differentially expressed in B. goodwinii 184 only and with A. biguttatus inoculations (Padj = 0.04, B. goodwinii inoculation only; Padj = 185 0.0001, B. goodwinii and A. biguttatus co-infection). The AvrE T3 effector is found in a wide 186 number of bacterial plant pathogens due to its proclivity for horizontal gene transfer [23]. 187 Notably, within the plant pathogen *Pseudomonas viridflava*, AvrE is the primary virulence 188 factor [24].

189

190 Detoxification genes in *B. goodwinii* are stimulated by the presence of *A. biguttatus*,
191 which may neutralise host defences

192 As described above, co-infection of oak logs with A. biguttatus significantly increases the 193 number of significantly differentially expressed genes within B. goodwinii and stimulates 194 expression of putative pathogen genes. In addition, homologs of genes which neutralise tree 195 defences were expressed. In previous studies, these homologs have been shown to create a 196 desirable environment for pupation and bacterial persistence [25]. The number of 197 significantly differentially expressed genes in *B. goodwinii* inoculated logs increased from 198 191 to 552 when A. biguttatus eggs were co-inoculated. Genes upregulated by A. biguttatus 199 eggs and not in B. goodwinii only log inoculations included host defence detoxification 200 genes; catalase peroxidase (Padj < 0.0001; E.C. 1.11.1.21), glutathione reductase (Padj =201 0.02; E.C. 1.8.1.7), and glutathione regulated potassium efflux system (Padj = 0.02). Catalase 202 peroxidase and glutathione reductase are encoded on the same operon; catalase peroxidase 203 (katG) protects against hydrogen peroxide released by host defences [26] and glutathione is a 204 metabolite of isoprene and its derivative terpene, both of which are common in oak trees and 205 used to combat abiotic stress and in high quantities are toxic to bark boring beetles [7,27,28]. 206 B. goodwinii mediated terpene reduction may exhaust terpene synthesis similar to that of 207 drought stressed oaks which initially produce abundant amounts of terpenes but upon severe 208 drought stress are no longer able to synthesise the volatiles, leaving them open to herbivores 209 [29].

210

The oak host up-regulates more defence-associated genes during co-inoculation with *A*. *biguttatus*

213 Examination of oak host transcripts within infection tests revealed differential gene 214 expression when challenged with B. goodwinii only compared to B. goodwinii with A. 215 biguttatus eggs. This analysis revealed 25 significantly up-regulated genes in logs inoculated 216 with B. goodwinii and A. biguttatus eggs compared to 12 up-regulated genes with only B. 217 goodwinii. This result provides futher evidence of an increase in activity of B. goodwinii 218 when co-infected with A. biguttatus. For both B. goodwinii treatments we discovered the up-219 regulation of genes encoding the calcium sensor protein CML38. This protein, and calcium 220 signalling proteins in general are reportedly induced during, wounding, stress and pathogen 221 infection [30,31]. Furthermore, during inoculation with B. goodwinii only, and with G. 222 quercinecans and eggs, there was significant up-regulation of a NDR1/HIN1 like protein, 223 which is associated with senescence and pathogen infection [32]. Host genes encoding 224 NDR1/HIN1 like proteins have previously been reported as up-regulated when comparing field AOD lesion bark to that from non-symptomatic trees. During inoculation of *B. goodwinii* and eggs, there was also significant up-regulation of two infection associated genes encoding WUN1, a wound induced protein, and EP3, an endochitinase associated with infection [33–35]. These results support the conclusion that bacterial co-infection with *A. biguttatus* enhances not only bacterial activity but also overall triggering of host defenceassociated genes.

231

In vitro analysis of the *B. goodwinii* and *G. quercinecans* transcriptome response to oak sapwood and phloem tissue

To gain greater understanding of interactions between two key bacteria within the AOD pathobiome, *in vitro* transcriptome assays were designed to measure gene expression changes of *B. goodwinii* and *G. quercinecans* in pure cultures and co-cultures containing oak phloem and sapwood (figure 3 and see methods for recipe). A key unanswered question in AOD pathology relates to the nature of pathobiome interactions between *B. goodwinii* and *G. quercinecans*, and whether they represent competitive or cooperative strategies.

240

Gene expression of *B. goodwinii* within phloem and sapwood *in vitro* cultures varies substantially between single inoculations and co-cultures

Gene expression analysis revealed that *B. goodwinii* has a substantial transcriptomic response to oak sapwood tissue two hours post inoculation, significantly differentially expressing 39 genes (P < 0.05; 35 upregulated and 4 downregulated) (figure 3a). Upregulated genes were mostly sugar transport/catabolism (n = 11) and general metabolism genes but also included an anti-bacterial gene, the type I secretion protein colicin V (attacks closely related bacteria) [36]. This effect is not found in oak phloem tissue (figure 3b), indicating that *B. goodwinii* is stimulated by glucose and xylose rich sapwood tissue which it can utilise as a sugar source.

In co-culture, two hours post inoculation with *G. quercinecans*, *B. goodwinii* significantly differentially expressed genes which were not expressed in axenic *B. goodwinii* culture (n =14 in phloem; n = 13 in sapwood) (figure 3i). This response was found in both oak phloem and sapwood tissue (figure 3i – 3l), with upregulated genes including those associated with sugar depolymerisation, which hydrolyse long chain sugar polymers such as α -Narabinofuranosidae (E.C. 3.2.1.55), bacterial α -L-rhamnosidase (E.C. 3.2.1.40), and β galactosidase (E.C. 3.2.1.23). These enzymes degrade plant tissue by breaking glycosidic linkages in the pectic polysaccharide, rhamnogalacturonan-II [37] and hemicellulose [38]. In sapwood at two hours post inoculation (figure 3a), flagellar motility genes (n = 2) were upregulated including the motility regulator *fliA* [39] indicating that sapwood and *G*. *quercinecans* stimulate the flagellar apparatus of *B. goodwinii*.

261

G. quercinecans has a substantial upregulation of genes towards oak phloem tissue but not sapwood

The environmental reservoir and ecological niche of *G. quercinecans* is unconfirmed. However, it is a robust bacterium that can survive in harsh environments [40] and is consistently isolated from AOD lesions where it may contribute to tissue necrosis [18]. Evidence provided here reveals that *G. quercinecans* can is differentially stimulated by oak phloem (figure 3f) and may assist *B. goodwinii* in colonising this environment by inducing expression of hitherto unexpressed genes (figure 3j & 3l).

270 Here, G. quercinecans significantly differentially expressed 42 genes in single inoculations 271 with phloem tissue at two hours post inoculation (32 upregulated and 10 downregulated) 272 (figure 3f). A large number of upregulated genes are involved in sugar catabolism/transport 273 (n = 10), but also upregulated were general metabolism genes, the type IV secretion system 274 (T4SS) component virB4 and a key PCWDE - rhamnogalacturonan lyase (E.C. 4.2.2.23). The 275 in vitro environment, containing oak phloem and sapwood, may mirror the environmental 276 habitat of G. quercinecans, which has previously been isolated from rotting wood and has 277 many saprophytic properties [18,40].

278

279 Sugar consumption by G. quercinecans in oak sapwood is stimulated by B. goodwinii

280 Compared to axenic growth of G. quercinecans in sapwood (figure 3e & 3g), co-culture with 281 B. goodwinii induced significant differential expression of 21 genes (14 upregulated and 7 282 downregulated) (figure 3i & 3k). Upregulated gene function included sugar 283 catabolism/transport (n = 5), iron transporters (n = 3) and two secondary PCWDEs (n = 2). It 284 was anticipated that co-culture could potentially induce expression of anti-bacterial effectors 285 but similar to B. goodwinii in phloem, G. quercinecans catabolises and transports sugars from sapwood when B. goodwinii is present (figure 3i - 3l). Despite the encoding of multiple 286 287 toxin-antitoxin systems and type VI secretion systems, there was no evidence of competitive 288 behaviour between B. goodwinii and G. quercinecans. These are closely related bacteria,

isolated from the same environmental niche and these experiments suggest that they assist
each other to metabolise oak tissue. Anti-bacterial effectors may be expressed at later stages
of co-culture, when resources are reduced, but this was not tested here.

292

RNA-seq validation using RT-qPCR analysis of *G. quercinecans* FRB97 and *B. goodwinii* FRB141 putative pathogenicity genes

Two RT-qPCR gene expression assays were used to validate RNA-seq data using the same RNA extracts as the *in vitro* RNA-seq experiment. In *G. quercinecans tssD* was selected, as homologs of this gene form part of the T6SS injectosome [41], and in *B. goodwinii fliA* was selected, which is an alternative sigma factor and controls flagella filament synthesis, chemotaxis machinery, and motor switch complex genes in *E. coli* [42].

300 RT-qPCR assays revealed that gene expression was highest at 6 HPI for *tssD* (an average of 2.2×10^5 absolute transcript copies at 2 HPI, 3.5×10^6 at 6 HPI, 2.7×10^5 at 12 HPI, 4.1×10^4 at 301 24 HPI), and 2 HPI for *fliA* (an average of 5.5x10⁴ absolute transcript copies at 2 HPI, 302 8.7×10^3 at 6 HPI, and 2.6×10^3 at 12 HPI) (electronic supplementary material figure, S1). 303 304 RNA-seq data revealed high gene expression of *fliA* in axenic *B*. goodwinii culture at 2 HPI, 305 and differential upregulation in co-culture with G. quercinecans, in nutrient broth (NB) & 306 sapwood (NBS) and nutrient broth & phloem (NBP) cultures at 2 HPI only, with gene 307 expression being suppressed with the addition of G. quercinecans in Nutrient Broth (NB). 308 tssD was highly expressed at 6 HPI, concurring with the RT-qPCR data (electronic 309 supplementary material figure, S1), and was differentially upregulated at 2 HPI in NBS and 310 NBP compared to NB. Within the G. quercinecans transcriptome tssD was upregulated in 311 NBS and NBP, suggesting that it is part of a wider virulence transcription cascade, and may 312 respond to eukaryotic stimuli. Transcriptomic expression data of *tssD* and *fliA*, data correlates 313 with the RT-qPCR data, however, small variations may be explained by the high sensitivity 314 of RT-qPCR [43,44].

315

317 Conclusions

Here we investigated the emergent properties of pathobiome virulence in AOD. We used 318 319 gene expression analysis of axenic and co-cultures of bacteria supplemented with oak inner 320 bark tissue, and oak infection tests using combinations of the A. biguttatus bettle and 321 microbial pathobionts. We demonstrated that the pathogenic potential of the dominant 322 bacterial species within the AOD lesion pathobiome, B. goodwinii, is stimulated by a co-323 invading native beetle, A. biguttatus, and also potentially induced by other microorganisms in 324 the AOD pathobiome associated with either the host or A. biguttatus. Furthermore, B. 325 goodwinii genes induced by the presence of A. biguttatus may confer nutrient acquisition 326 benefits to beetle eggs and larvae.

327 The co-operative behaviour of B. goodwinii and G. quercinecans in a nutrient rich 328 environment may differ from the AOD lesion environment where resources are scarce. 329 However, both bacteria persisted in oak phloem and sapwood when combined, and when 330 resources were plentiful there was no significant upregulation of interbacterial competition 331 genes. It was also revealed that G. quercinecans favours sugar metabolites from oak phloem 332 tissue, whereas B. goodwinii favours oak sapwood as a carbon source. The role of L. 333 britannica in the lesion environment is unclear but merits further investigation due to its 334 encoded pathogenic potential and high expression activity in combination with B. goodwinii 335 and A. biguttatus. It is possible that AOD pathobiome constituents each contribute 336 degradative enzymes to systematically macerate oak tissue, thereby co-operating to provide 337 ingestible sugars as a public good. To fully characterise the molecular processes uncovered in 338 this study will require tractable genetic manipulations of single gene effects in appropriate 339 model systems.

In conclusion, we identified expression of key pathogenicity genes in *Brenneria goodwinii*, the dominant member of the AOD pathobiome, tissue-specific gene expression profiles, cooperation with other bacterial pathobiome members in sugar catabolism, and demonstrated amplification of pathogenic gene expression in the presence of *Agrilus* larvae. These data highlight the emergent propereties of complex multidimensional interactions between host plants, insects and the microbiome that underpin complex tree decline diseases that threaten the global landscape.

- 348

347

349 In vitro culture-based assay

Methods

350 Strains, growth medium and conditions

351 Strains of Gibbsiella quercinecans FRB97 and Brenneria goodwinii FRB141 were obtained 352 by Forest Research (Surrey, UK) from AOD affected trees. Isolates were maintained on 353 nutrient agar (Oxoid) at room temperature. To simulate growth on sapwood and phloem, cells 354 were cultured in nutrient broth (Oxoid) containing 1% (w/v) milled sapwood (NBS), nutrient 355 broth with 1% (w/v) milled phloem (NBP) and a control consisting of nutrient broth (NB). 356 Initially, a 10 ml starter culture from a single colony was incubated overnight to stationary 357 phase at 28°C on a shaking incubator at 100 rpm. 1% of the overnight culture, was 358 centrifuged and re-suspended, before addition to three replicate culture flasks containing 150 359 ml volumes of NB, NBS, and NBP (figure 3). The flasks were incubated at 28°C and 100 360 rpm, for 6 HPI, with cell suspensions collected at 2 HPI and 6 HPI. At each time point 25 ml 361 of liquid was collected in a 50 ml Falcon tube and centrifuged for 5 mins at 3000 rpm. The 362 supernatant was discarded, and pelleted cells were frozen in liquid nitrogen.

363

364 Log infection assay

365 Log trials were established in 2015 (electronic supplementary table, S3 for list of log 366 inoculation treatments, resultant lesion sizes and futher information). Owing to the high cost 367 of transcriptomics when the trial was terminated and samples processed, only a sub-set of 3 368 inoculations points in each of the above described treatments were sampled, at random, from 369 the log test, except where there were exceptional cases of typical AOD lesions i.e. two 370 Lonsdalea inoculations, which were specifically included in the transcriptomic analyses. 371 Following lesion area measurements and plating lesion margin wood chips [8] the remaining 372 lesion was chiselled out, placed in a labelled ziplock plastic bag and snap-frozen in liquid 373 nitrogen and stored at -80C until RNA extraction too place.

374

375 **RNA extraction**

376 RNA extraction from bacterial cultures

377 Total RNA was extracted from cell pellets of bacterial cultures using the RNeasy Mini Kit (Qiagen), according to manufacturer's instructions. Genomic DNA was removed from 378 379 extracted RNA samples using TURBO DNA-free DNase kit (Ambion). Total RNA was 380 pooled from three biological replicates in equimolar quantities giving a total quantity of 750 381 ng (electronic supplementary material figure, S2). Total rRNA was depleted to enrich mRNA 382 (transcripts) using the RiboZero rRNA depletion kit (Illumina). The protocol was performed 383 according to manufacturer's instructions. Depleted mRNA concentrations were measured 384 using a Qubit fluorometer (Invitrogen). Remnant rRNA was minimal as confirmed by the 385 Centre for Genomic Research (CGR) (University of Liverpool, UK), using the Agilent 2100 386 BioAnalyzer.

387

388 RNA extraction from log inoculations

389 RNA was extracted from logs using the method described in our previous multi-omic AOD 390 work and described here [45]. Briefly, inner bark around log inoculation spots was scraped 391 off and snap frozen in liquid nitrogen. Oak tissue was homogenised using a mortar and pestle, 392 and extraction buffer (4 M guanidine thiocyanate, 0.2 M sodium acetate pH 5.0, 25 mM 393 EDTA, 2.5% (w/v) polyvinylpyrrolidone and 1% (v/v) β -mercaptoethanol) was added. The 394 frozen tissue in extraction buffer was further ground until thawed, while additional extraction 395 buffer and 20% sodium lauroyl sarcosinate were mixed into the sample. The sample mixture 396 was shaken vigorously at room temperature and further processed using the RNeasy Plant 397 Mini kit (Qiagen). After centrifugation in the QIAShredder column, 350 µl of the supernatant 398 was mixed with 0.9 volumes of ethanol, and subsequently centrifuged in the RNeasy Mini 399 column. After this centrifugation step, the manufacturer's instructions for the RNeasy Plant 400 Mini kit were followed. The extracted RNA was treated with DNase I (Qiagen) and further 401 concentrated and purified using the RNeasy MinElute Cleanup kit (Qiagen) following the 402 manufacturer's instructions. The purified RNA was checked for quality using 1% agarose gel 403 electrophoresis and a NanoDrop spectrophotometer (LabTech), and the concentration 404 determined using the Qubit RNA HS assay kit (Thermo Fisher) following the manufacturer's 405 instructions. Subsequently, rRNA was depleted from RNA extracts using a 1:1 combination 406 of the Ribo-Zero rRNA Removal kits for plant seed/root and for bacteria (Illumina) according 407 to the manufacturer's instructions. The rRNA depleted samples were again purified using the 408 RNeasy MinElute Cleanup kit (Qiagen) again and stored at -80 °C before sequencing.

410 **RNA sequencing**

411 Library preparation, transcriptomic sequencing, and post-sequencing OC of depleted RNA 412 samples was performed by Centre for Genomic Research (CGR), University of Liverpool, 413 UK. Samples were assayed for quality using an Agilent 2100 Bioanalyzer. Log infection 414 samples were further assayed for quality using the Eukaryote Total RNA Pico Series II. All 415 libraries were prepared using the strand-specific ScriptSeq kit (Illumina), and subsequently 416 paired-end sequenced (2x125 bp) on one lane (N.B. in vitro and log infection samples were 417 sequenced on separate lanes) of the Illumina HiSeq 2500 platform (electronic supplementary 418 material figure, S3 & electronic supplementary material figure, S4).

419

420 Transcriptome analysis

421 RNA-seq QC

Illumina adapter sequences were removed from raw FastQ files containing the sequencing reads using Cutadapt v1.2.1 [46], using the option –O 3, which specifies that at least 3 base pairs have to match the adapter sequences before they were trimmed. Sequences were quality trimmed using Sickle v1.2 [47] with a minimum quality score of 20. Reads shorter than 10 bp were removed. RNA-seq QC was performed by Centre for Genomic Research (CGR), University of Liverpool, UK (electronic supplementary material figure, S3 & electronic supplementary material figure, S4).

429

430 Bioinformatic analysis of transcriptome data

431 Bioinformatic analyses were carried out on SuperComputing Wales, an HPC network, using 432 GNU/Linux Red Hat Enterprise Linux Server release 7.4 (Maipo). A complete list of 433 commands used to perform the below analysis is hosted on GitHub 434 (https://github.com/clydeandforth/Bg_Ab_logs.git).

435

436 Transcriptome alignment and differential gene expression analysis

RNA recovered from log inoculations and sequenced on the Illumina HiSeq, was aligned
using Bowtie2 v1.1.2 [48] to an in-house database of structurally and functionally annotated
coding regions (electronic supplementary methods) used in a previous field AOD

440 microbiome analysis [15], but with the addition of Lonsdalea britannica 477. Transcript 441 counts for each gene were calculated using eXpress v1.5.1 [49]. To give an overview of 442 species activity in the lesion environment, an active gene was defined as those with 443 transcripts per million (TPM) >1 and a total transcript count of three. TPM rather than raw 444 read counts was used to normalise the number of transcripts across samples and remove 445 sequencing depth as an experimental artefact. Subsequently, in a separate test, to get a 446 statistically robust understanding of transcriptional activity, significantly differentially 447 expressed genes were identified using DESeq2 v1.2 [50]. Genes which had P-adjusted values 448 <0.05 between conditions were considered as significantly differentially expressed. Principal 449 coordinate analyses based on dispersion of mean normalised gene count data between 450 samples was calculated and plotted using DESeq2 v1.2.

451 Gene-set enrichment analyses of KEGG pathways were used to measure functional 452 upregulation of gene families between samples using the R packages gage v2.30.0 [51] and 453 clusterProfiler v3.8.1 [52]. GAGE uses a two sample t-test to compare expression level 454 changes between genesets. KEGG datasets were compiled from KEGGREST v1.20.1 455 (accessed 04/02/2019) and comprised pathways from the following bacteria: Dickeya 456 dadantii 3937, Pectobacterium carotovorum subsp. carotovorum PC1, Escherichia coli K12, 457 E. coli 0157:H7 Sakai, Rahnella aquatilis CIP 78.65, Serratia proteamaculans 568, and 458 plants: Phoenix datylifera, Arabidopsis thaliana, Methylorubrum populi BJ001.

459

460 Multivariate analysis - Generalised linear model

To test for biological variation between samples, the effect of inoculum, beetle presence/absence, lesion size and replicate were included in a generalised linear model (GLM) [53]. Normalised read count data produced using eXpress (described below) were set as the multivariate response variable and the biological predictors were fit using a negative binomial distribution. The 'manyglm' function of the R package [54] mvabund [55] was used to carry out the analysis. Inoculum, beetle presence/absence, lesion size, and replicate were included as exploratory variables to allow the model to test our hypotheses.

468

469 Transcriptomic analysis of *in vitro* sequence data

470 Sequenced RNA from *in vitro* tests was aligned to a custom database and counted as 471 described above. The transcript per million (TPM) counts from eXpress analysis were used to

- 472 calculate the Generalised Fold Change (GFOLD) [56], which uses the posterior distribution
- 473 of the raw fold change to calculate differential expression of genes between conditions and is
- 474 analogous to the *P* value in DESeq2. Genes which had GFOLD values >1.5 or <1.5 between
- 475 conditions were considered as significantly differentially expressed.
- 476
- 477 Data availability
- 478 Sequence data has been deposited in NCBI under BioProject PRJNA369790.
- 479

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- 488

489 **Conflict of interest**

- 490 The authors declare no conflicts of interest.
- 491

492 Author contributions

493 JD and MB carried out the molecular lab work, RNA extraction and depletion, statistical 494 and bioinformatic analysis. JD drafted the manuscript and created the figures; JEM 495 supervised the labwork and critically revised the manuscript; SD conducted log tests and 496 critically revised the manuscript; All authors, designed and coordinated the study. All 497 authors gave final approval for publication and agree to be held accountable for the work 498 performed therein.

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653 Figure legends

654 Figure I. Transcriptome analysis of oak log infection tests comprising single bacterial species 655 incoluations and bacteria plus Agrilus biguttatus egg inoculations. From left to right: Organisms 656 inoculated into oak logs are shown in the key on the top right, these are Gibbsiella quercinecans, 657 Brenneria goodwinii, Lonsdalea britannica and Agrilus biguttatus. There were three biological replicates of 658 each infection test, including replicate water only and wound controls. Each of the three bacterial 659 species were inoculated individually and in combination with eggs of A. biguttatus. Exemplary pictures 660 of a single log inoculation replicate from each treatment are shown. The number of expressed genes 661 from log inoculations are shown in the bar chart, with each expressed gene aligned against a custom 662 database and colour coded with the Genus/Species key shown on the bottom left of the figure. Oak 663 transcripts were excluded from the bar chart.

664

665 Figure 2. Transcriptome analysis of Brenneria goodwinii inoculations on live oak logs. (b) Gene set 666 enrichment analysis (GSEA) of B. goodwinii gene families when compared to (left) water and wound 667 control oak logs; (right) B. goodwinii inoculated in combination with A. biguttatus when compared to 668 B. goodwinii only. The lower q-value represents increased magnitude of gene family expression and 669 circle size represents number of genes per family. (a), principal component analysis (PCA) of (top) B. 670 goodwinii (n = 3) compared to control (n = 6); (bottom) B. goodwinii and A. biguttatus compared to 671 control (n = 6). (c) gene expression changes of selected significantly differentially expressed genes, 672 these are anti-toxicity genes (yellow), biofilm and persistence genes (purple), secretion system 673 effectors (blue). (top) B. goodwinii compared to control; (bottom) B. goodwinii and A. biguttatus 674 compared to control. Transcriptome samples were taken from log inoculations of bacterial 675 combinations, wound and water controls, and field samples of AOD lesions and asymptomatic oaks. 676 Bg = Brenneria goodwinii; Gq = Gibbsiella quercinecans; eggs = A. biguttatus.

677

678 Figure 3. In vitro transcriptome analysis of Brenneria goodwinii and Gibbsiella quercinecans in nutrient 679 broth supplemented with oak phloem and oak sapwood. Each panel shows gene expression changes 680 when phloem and sapwood are present, compared with nutrient broth only controls. (a) B. goodwinii 681 in sapwood at 2 HPI. (b) B. goodwinii in phloem at 2 Hours Post Inoculation HPI. (c) B. goodwinii in 682 sapwood at 6 HPI. (d) B. goodwinii in phloem at 6 HPI. (d) G. quercinecans in oak sapwood at 2 hours 683 post inoculation (HPI). (e) G. guercinecans in oak phloem at 2 HPI. (f) G. guercinecans in sapwood at 6 684 HPI. (g) G. quercinecans in phloem at 6 HPI. (i) B. goodwinii and G. quercinecans in sapwood at 2 HPI. 685 (j) B. goodwinii and G. quercinecans in phloem at 2 HPI. (k) B. goodwinii and G. quercinecans in sapwood

686	at 6 H	HPI. (I) B. goodw	/inii and (G. quercinecans i	in phloem at	t 6 HF	Pl. DEG	= differentially	expres	sed gene.
687	HPI	=	hours	post	inoculation.	GFOLD	is	the	generalised	fold	change.