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Genomic analyses confirm close relatedness between *Rhodococcus defluvii* and *Rhodococcus equi (Rhodococcus hoagii)*

Vartul Sangal¹*, Amanda L. Jones¹, Michael Goodfellow², Paul A. Hoskisson³, Peter Kämpfer⁴, Iain C. Sutcliffe¹

¹Faculty of Health and Life Sciences, Northumbria University, Newcastle upon Tyne NE1 8ST, UK

²School of Biology, University of Newcastle, Newcastle upon Tyne NE1 7RU, UK

³Strathclyde Institute of Pharmacy and Biomedical Sciences, University of Strathclyde, 161 Cathedral Street, Glasgow G4 0RE, UK

⁴Institut für Angewandte Mikrobiologie, Justus-Liebig-Universität, Giessen, D-35392, Germany

*Correspondence: Vartul Sangal, Faculty of Health and Life Sciences, Northumbria University, Northumberland Building, Newcastle upon Tyne – NE1 8ST, UK. Tel: +44 191 243 7173; e-mail: <u>vartul.sangal@northumbria.ac.uk</u>

Keywords: *Rhodococcus equi*, *Rhodococcus defluvi*, genome, average nucleotide identity, average amino-acid identity

1 Abstract

8

Rhodococcus defluvii strain Ca11^T was isolated from a bioreactor involved in extensive
phosphorus removal. We have sequenced the whole genome of this strain and our
comparative genomic and phylogenetic analyses confirm its close relatedness with *Rhodococcus equi (Rhodococcus hoagii)* strains, which share >80% of the gene content. The *R. equi* virulence plasmid is absent though most of the chromosomal *R. equi* virulenceassociated genes are present in *R. defluvii* Ca11^T. These data suggest that although *R. defluvii*

is an environmental organism, it has the potential to colonise animal hosts.

9 Rhodococcus defluvii is a Gram-positive, mycolic acid-containing, rod shaped actinobacterium that has been described as a new member of the heterogeneous genus 10 Rhodococcus (Jones and Goodfellow 2012; Kämpfer et al. 2014). The type strain of this 11 species, Ca11^T (=DSM 45893^T =LMG27563^T), was isolated from a wastewater treatment 12 bioreactor involved in phosphorus removal. Strain Ca11^T showed the highest 16S rRNA 13 sequence similarity (98.9%) and corresponding DNA-DNA relatedness value (51.3%; 14 reciprocal 38.1%) to the type strain of Rhodococcus equi (Rhodococcus hoagii; Kämpfer et 15 al., 2014). The nomenclature of these taxa is currently a matter of debate as the priority of the 16 17 name R. hoagii over R. equi (or vice versa) is under review by the Judicial Commission of the International Committee on Systematics of Prokaryotes (Garrity 2014) while the bacterial 18 genus name Rhodococcus is considered to be illegitimate (Tindall 2014). For clarity, we here 19 20 refer to the R. equi/R. hoagii taxon as R. equi. In this study, we have sequenced the genome of *R*. *defluvii* strain $Ca11^{T}$ and 21 performed comparative analyses with the genome sequences of *R. equi* strains $C7^{T}$ (Sangal et 22 23 al. 2014), 103S (Letek et al. 2010) and ATCC 33707 (Qin et al. 2010) [GenBank accession numbers APJC00000000, NC_014659 and NZ_CM001149, respectively]. Genomic DNA 24 extracted from 1.5ml of culture grown for 48 h at 30°C in Brain-Heart Infusion broth (Oxoid) 25 was sequenced on an Illumina MiSeq instrument, according to the manufacturer's 26 27 instructions. A total of 2,156,061 reads with an average read length of 238 bp were assembled 28 into 267 contigs (>200 bp) using CLC Genomic Workbench (Qiagen). The size of assembly was 5,134,337 bp with an average 75-fold coverage. 29 The size of the draft genome and G+C content of *R*. *defluvii* strain Ca11^T (5.13 Mb, 30 68.71%) are similar to those of *R. equi* strains $C7^{T}$ (5.20 Mb, 68.79%), 103S (5.04 Mb, 31 68.82%) and ATCC 33707 (5.26 Mb, 68.77%). However, the genome sequence has only 32 been completed for strain 103S and so these values may slightly vary for other strains if their 33

genomes are finished. Using the RAST pipeline (Aziz et al. 2008), the Ca11^T genome was 34 annotated to have 4,796 features including 4,740 protein coding sequences. The genomes of 35 R. equi strains were also re-annotated using the RAST pipeline to allow an equivalence of 36 annotation. The Call^T genome was found to share 4.166 genes with the three R. equi strains 37 (3,720 with bi-directional and 446 with uni-directional protein BLAST hits; Aziz et al. 2012). 38 It also shared an additional 128 genes with at least one R. equi strain but not with all three. 39 446 genes were specific to *R. defluvii* Cal1^T that were absent in the *R. equi* genomes; 361 of 40 these encode hypothetical proteins and six belong to mobile genetic elements (transposase, 41 42 phage associated or mobile element proteins). A BLAST search of 75 randomly selected hypothetical proteins of *R. defluvii* against the NCBI protein database using default settings 43 44 revealed homologies for most of them with hypothetical proteins in other rhodococci or other bacterial species (data not shown), indicating that not all are unique to *R. defluvii* Ca11^T. The 45 remaining 79 genes specific to R. defluvii Call^T (compared to the R. equi strains) can 46 typically be related to known metabolic activities (Table S1), including a gene encoding 47 alkylphosphonate utilization protein PhnA. The *phn* operon gene products are involved in the 48 cleavage of carbon-phosphorus bonds in alkylphosponates (Chen et al. 1990). However, the 49 presence of the *phnA* gene in strain Ca11^T is unlikely to be associated with phosphorus 50 removal in the bioreactor from which it was isolated because the other genes of this operon 51 are missing. Three homologs of *phnB* and two homologs of *phnE* genes were present 52 elsewhere in the Call^T genome but they are shared with the *R. equi* strains. A number of 53 54 other genes involved in phosphorus metabolism are also common between R. defluvii and the three R. equi strains. 55

An operon in the genome of strain Ca11^T that encodes Ter family proteins (TerA, TerB, TerC-like and two TerD) and associated biosynthetic enzymes is absent from the genomes of the three *R. equi* strains (Table S1). Comparable loci have previously been

59 suggested to be involved in biosynthesis of nucleoside-like metabolites (Anantharaman et al. 2012). The protein BLAST search revealed the presence of homologs of these genes in other 60 rhodococci and actinomycetes, suggesting a potential horizontal acquisition of this operon by 61 62 R. defluvii. Alternatively, this operon may have been lost by R. equi as it has adapted to a pathogenic lifestyle. Two of the genes specific to *R. defluvii* Call^T (compared to the *R. equi* 63 64 strains) encode phospholipase C enzymes. Phospholipases C are the virulence factors that induce alveolar macrophage necrosis, resulting in cell death (Assis et al. 2014). As noted 65 above, most of the genes specific to strain Ca11^T encode hypothetical proteins and it is 66 67 possible that some of these uncharacterized proteins contribute to functional variations between R. defluvii and R. equi. 68

Rhodococci are generally involved in environmental processes such as the 69 70 degradation of organic and xenobiotic substances, except for the pathogens R. equi and Rhodococcus fascians (Bell et al. 1998; Alvarez 2010). The pathogenicity of these two 71 72 species has been associated with the presence of large plasmids encoding virulence proteins 73 (Takai et al. 2000; Letek et al. 2008; Francis et al. 2012; Stes et al. 2013). The virulence plasmid in R. equi is 80-90 Kb in size and carries a pathogenicity island encoding virulence 74 associated proteins (Vap) while plasmid free strains were found to be avirulent (Takai et al. 75 2000). A sequence BLAST-based functional comparison using the SEED server (Aziz et al. 76 2012) revealed the absence of Vap proteins (VapA, C-I proteins from plasmid pVAPA1037 77 and VapB, J-M from pVAPB1593; Letek et al. 2008) in the draft genome sequence of R. 78 *defluvii*, suggesting the absence of the virulence plasmid in strain Ca11^T. However, 228 of the 79 243 R. equi chromosomal virulence-related genes defined by Letek et al. (2010) are present 80 in strain Call^T (Table S2), including the *esx* cluster. The *paa* operon that was identified in *R*. 81 equi strain ATCC 33707 and which may be involved in pathogenesis in humans (Sangal et al. 82 2014) is absent from *R. defluvii* strain Ca11^T. The presence of a high proportion of virulence-83

related genes in the genome of strain Ca11^T suggests that this organism may also have the
potential to colonise animal hosts. Indeed, it is noted that three additional bacterial strains
with 16S rRNA gene sequences identical to that of strain Ca11^T have been isolated from
salmon intestines (Skrodenyte-Arbaciauskiene,V. & Virbickas T. Genbank accession
numbers HM244990, HM244992 and HM244993).

A phylogenetic analysis was performed using PhyloPhlAn (Segata et al. 2013) 89 including Rhodococcus erythropolis PR4 (Sekine et al. 2006), Rhodococcus jostii RHA1 90 (McLeod et al. 2006). Nocardia brasiliensis ATCC 700358 (Vera-Cabrera et al. 2012) and 91 Corynebacterium diphtheriae NCTC 05011 (Sangal et al. 2012) were used as outgroups. 92 PhyloPhlAn automatically extracts the sequences of the 400 most conserved universal 93 94 proteins that were identified by off-line pre-processing of all available microbial genomes by 95 Segata et al.(2013). It generates highly robust phylogenetic trees from a concatenated alignment of computationally selected subset of amino-acid sequences with highest entropy 96 97 and an appropriate relative contribution of the most conserved residues from each protein 98 following a maximum likelihood maximization approach (gamma model of rate heterogeneity) with 20 bootstrap replicates using RAxML (Stamatakis 2006). Our 99 PhyloPhlAn analysis showed that *R. defluvii* Ca11^T shared a phyletic line with *R. equi* that 100 was relatively distant from the other rhodococci and from N. brasiliensis (Fig. 1). BLAST-101 based average nucleotide identities (ANIb) between the genomes of *R*. *defluvii* Ca11^T and the 102 R. equi strains were 82.96-83.25% (Richter and Rosselló-Móra 2009) and average amino acid 103 identities (AAI) varied between 85.31-85.45%. The ANIb and AAI values between R. 104 defluvii and the other rhodococci (R. jostii RHA1 and R. erythropolis PR4) were < 76% and 105 <72%, respectively. The digital DNA-DNA hybridization (dDDH) distances were calculated 106 using the genome-to-genome distance calculator at the GGDC 2.0 web server (Auch et al. 107 2010; Meier-Kolthoff et al. 2013). GGDC values mimic conventional DNA-DNA 108

| 109 | hybridization values and have been shown to have very high correlation with 16S rRNA |
|-----|---|
| 110 | sequence distances (Auch et al. 2010; Meier-Kolthoff et al. 2013). GGDC 2.0 uses three |
| 111 | different formulae to calculate the distances and the results of formula-2, which has been |
| 112 | recommended for analysing draft genomes (Auch et al. 2010), were considered in this study. |
| 113 | The dDDH values between <i>R. defluvii</i> and <i>R. equi</i> strains C7 ^T , 103S and ATCC 33707 were |
| 114 | 26.9 ± 3.02 , 27 ± 3.02 and 27.1 ± 3.01 , respectively. The <i>R. defluvii</i> genome showed lower |
| 115 | dDDH similarities with the <i>R. erythropolis</i> PR4 (20.2 \pm 2.73) and <i>R. jostii</i> RHA1 (20.7 \pm |
| 116 | 2.81) genomes, values that are comparable to the dDDH distances from <i>N. brasiliensis</i> ATCC |
| 117 | 00358 (20.4 \pm 2.63) and <i>C. diphtheriae</i> NCTC 05011 (21 \pm 2.53). Cumulatively, these results |
| 118 | suggest that R. defluvii is more closely related to R. equi than to other rhodococci, as |
| 119 | previously concluded from 16S rRNA gene sequence analysis (Kämpfer et al. 2014). |
| 120 | In addition to the nomenclatural issues highlighted above, it has been proposed that R . |
| 121 | equi should be reclassified as 'Prescottella equi' (Jones et al. 2013b; Jones et al. 2013a). |
| 122 | However, the genus name 'Prescottella' cannot be validated until the Judicial Commission |
| 123 | reports on whether the species epithet equi should be conserved over hoagii (Garrity 2014). |
| 124 | Based on the phylogenetic and genomic distances between <i>R. defluvii</i> and the other |
| 125 | rhodococci (Fig. 1), R. defluvii could eventually be reclassified as a second species within |
| 126 | 'Prescottella'. However, this conclusion needs further support from analyses of a larger |
| 127 | collection of genomes of <i>Rhodococcus</i> species. |
| 128 | In summary, we report the genome sequence of the type strain of the recently |
| 129 | identified species, <i>R. defluvii</i> strain Ca11 ^{T} . The strain is phylogenetically closely related to <i>R</i> . |
| 130 | equi strains with high similarities both at the nucleotide and functional levels. The whole |
| 131 | genome shotgun sequence has been deposited at DDBJ/EMBL/GenBank under the Accession |
| 132 | number JPOC00000000. The version described in this study is the first version, |
| 133 | JPOC01000000. |

| 134 | |
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| 140 | |

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227 Figure Legend

- **Figure 1.** Phylogenetic tree (radial, un-rooted) derived from 400 universal proteins using the
- program PhyloPhlAn showing the relatedness of *R. defluvii* Ca11^T with *R. equi* and
- 230 representatives of other closely related taxa. Scale bar shows normalized fraction of total
- branch lengths as described by Segata et al. (2013).

