Improved genome of *Agrobacterium radiobacter* type strain provides new taxonomic insight into *Agrobacterium* genospecies 4

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**ABSTRACT**

The reported *Agrobacterium radiobacter* DSM30174ᵀ genome is highly fragmented, hindering robust comparative genomics and genome-based taxonomic analysis. We re-sequenced the *Agrobacterium radiobacter* type strain, generating a dramatically improved genome with high contiguity. In addition, we sequenced the genome of *Agrobacterium tumefaciens* B6ᵀ, enabling for the first time, a proper comparative genomics of these contentious *Agrobacterium* species. We provide concrete evidence that the previously reported *A. radiobacter* type strain genome (Accession Number: ASXY01) is contaminated which explains its abnormally large genome size and fragmented assembly. We propose that *Agrobacterium tumefaciens* be reclassified as *A. radiobacter* subsp. *tumefaciens* and that *A. radiobacter* retains it species status with the proposed name of *A. radiobacter* subsp. *radiobacter*. This proposal is based, first on the high pairwise genome-scale average nucleotide identity supporting the amalgamation of both *A. radiobacter* and *A. tumefaciens* into a single species. Second, maximum likelihood tree construction based on the concatenated alignment of shared genes (core genes) among related strains indicates that *A. radiobacter* NCPPB3001 is sufficiently divergent from *A. tumefaciens* to propose two independent sub-clades. Third, *A. tumefaciens* demonstrates the genomic potential to synthesize the L configuration of fucose in its lipid polysaccharide, fostering its ability to colonize plant cells more effectively than *A. radiobacter*.

**INTRODUCTION**
The taxonomy and phylogeny of the genus *Agrobacterium* has proven to be complex and controversial. Bacteria of the genus *Agrobacterium* have been grouped into six species based on the disease phenotype associated, in part, with the resident disease-inducing plasmid. Among those six species are *A. tumefaciens* causing crown gall on dicotyledonous plants, stone fruit and nut trees and *A. radiobacter* that is not known to cause plant diseases of any kind (Bouzar & Jones 2001; Conn 1942; Kerr & Panagopoulos 1977; Panagopoulos et al. 1978; Riker et al. 1930; Starr & Weiss 1943; Süle 1978). An alternative classification approach grouped *Agrobacterium* organisms into three biovars based on physiological and biochemical properties without consideration of disease phenotype (Keane et al. 1970; Kerr & Panagopoulos 1977; Panagopoulos et al. 1978). The species and biovar classification schemes do not coincide well, in a large part, because of the disease-inducing plasmids, tumor-inducing (pTi) and hairy root-inducing (pRi), are readily transmissible plasmids (Young et al. 2001).

Many widely used approaches for bacterial species definition include composition of peptidoglycan, base composition of DNA, fatty acid and 16S rDNA sequence (Stackebrandt et al. 2002) in addition to newer methods based on the whole-genome analysis (Coutinho et al. 2016; Jain et al. 2017), horizontal gene transfer analysis (Bobay & Ochman 2017) or the core genome analysis (Moldovan & Gelfand 2018) which is used in the present study. The genus *Agrobacterium* is a prime example with many proposals and oppositions regarding the amalgamation of *Agrobacterium* and *Rhizobium* over the last three or four decades (Farrand et al. 2003; Gaunt et al. 2001; Young et al. 2001; Young et al. 2003). However, more recent studies appear to favor the preservation of the genus *Agrobacterium* backed by strong genetic and genomic evidence (Gan & Savka 2018; Ramírez-Bahena et al. 2014). Within the genus *Agrobacterium*, the taxonomic status of *A. radiobacter* and *A. tumefaciens* remains contentious (Sawada et al. 1993; Young 2008; Young et al. 2006). *Agrobacterium radiobacter* (originally proposed as *Bacillus radiobacter*) is a non-pathogenic soil bacterium associated with nitrogen utilization isolated more than a century ago in 1902 (Beijerinck & van Delden 1902; Conn 1942). On the other hand, *A. tumefaciens* (previously *Bacterium tumefaciens*) is a plant pathogen capable of inducing tumorigenesis (Smith & Townsend 1907). However, the descriptive assignment for *A. tumefaciens* was later found to be contributed by a set of genes located on the large Ti plasmid that can be lost (Gordon & Christie 2014). In other words, the curing of Ti plasmid in *A. tumefaciens* will change its identity to the non-pathogenic species, *A. radiobacter*. Furthermore, comparative molecular analysis based on single-copy housekeeping genes also supports the close relatedness of *A. radiobacter* and *A. tumefaciens*, blurring the taxonomic boundaries between these species (Mousavi et al. 2015; Shams et al. 2013). As taxa are reclassified into different populations that do not conform to the characteristics of the original description, the given names lose their significant and descriptive importance. Consistent with the Judicial Commission according to the Rules of the International Code of Nomenclature of Bacteria, Tindall (2014) concluded that “the combination of *A. radiobacter* has priority over the combination *A. tumefaciens* when the two are treated as members of the same species.
since \textit{A. radiobacter} was the first proposed and described in 1902 whereas \textit{A. tumefaciens} was first proposed and described in 1907) (Tindall 2014). However, given that \textit{A. tumefaciens} has been more widely studied than \textit{A. radiobacter} due to its strong relevance to agriculture (Bourras et al. 2015), it remains unclear but interesting to see if the broader scientific community will obey this rule by adopting the recommended species name change in future studies.

To our knowledge, a detailed comparative genomics analysis of \textit{A. radiobacter} and \textit{A. tumefaciens} type strains has not been reported despite their genome availability (Zhang et al. 2014). The high genomic relatedness of both type strains was briefly mentioned by Kim and Gan (2017) through whole genome alignment and pairwise nucleotide identity calculation from homologous regions. However, evidence is now mounting that the \textit{A. radiobacter} DSM 30147\textsuperscript{T} reported by Zhang et al. (2014) is contaminated, warranting immediate investigation (Jeong et al. 2016). The assembled genome is nearly 7 MB, the largest among \textit{Agrobacterium} currently sequenced at that time with up to 6,853 predicted protein-coding genes contained in over 600 contigs. At sequencing depth of nearly 200\times, its genome assembly is unusually fragmented even for a challenging microbial genome (Utturkar et al. 2017). Furthermore, the phylogenomic placement of \textit{A. radiobacter} DSM 30147\textsuperscript{T} based on this genome assembly has been questionable as evidenced by its basal position and substantially longer branch length relative to other members of the species (Gan & Savka 2018). The overly fragmented nature of this assembly also precludes fruitful comparative genomics focusing on gene synteny analysis. More importantly, analysis done on a contaminated assembly but with the assumption that it is not, will likely lead to incorrect biological interpretations (Allnutt et al. 2018).

In this study, we sequenced the whole genome of \textit{A. radiobacter} using a type strain that was sourced from the National Collection of Plant Pathogenic Bacteria (NCPBP). We produced a contiguous genome assembly exhibiting genomic statistics that are more similar to other assembled \textit{Agrobacterium} genomes. We show here, through comparative genomics and phylogenetics, that the previously assembled \textit{A. radiobacter} DSM 30147\textsuperscript{T} genome contains substantial genomic representation from another \textit{Agrobacterium} sp. isolated and sequenced by the same lab, consistent with our initial suspicion of strain contamination. Using the newly assembled genome for subsequent comparative analysis, we provide genomic evidence that \textit{A. radiobacter} DSM 30147\textsuperscript{T} and \textit{A. tumefaciens} B6\textsuperscript{T} are the same species. However, strain DSM 30147\textsuperscript{T} should not be considered as a merely non-tumorigenic strain of \textit{A. tumefaciens} as substantial genomic variation exists between these two type strains notably in the nucleotide sugar metabolism pathway that may contribute to their ecological niche differentiation.

\section*{MATERIALS & METHODS}

\subsection*{DNA extraction and whole genome sequencing}
Approximately 10 bacterial colonies were scrapped using a sterile P200 pipette tip from a 3-day-old nutrient agar culture and resuspended in lysis buffer with proteinase K.
(Sokolov 2000) followed by incubation at 56 °C for 3 hours. DNA purification was performed as previously described. The extracted DNA was normalized to 0.2 ng/μL and prepared using the Nextera XT library preparation kit (Illumina, San Diego, CA) according to the manufacturer’s instructions. The library was sequenced on an Illumina MiSeq desktop sequencer located at the Monash University Malaysia Genomics Facility (2 × 250 bp run configuration) that routinely sequences mostly decapod crustacean mitogenomes (Gan et al. 2016a; Gan et al. 2016b; Tan et al. 2015) and occasionally microbial genomes (Gan et al. 2015; Gan et al. 2014; Wong et al. 2014) without prior history of processing any member from the Agrobacterium genomospecies 4.

De novo assembly and genome completeness assessment
Raw paired-end reads were adapter-trimmed using Trimmomatic v0.36 (Bolger et al. 2014) followed by error-correction and de novo assembly using Spades Assembler v3.9 (Bankevich et al. 2012) (See Data S1 for specific trimming and assembly settings). Genome completeness was assessed with BUSCOv3 (Rhizobiales database) (Waterhouse et al. 2017).

Protein clustering
Gene prediction used Prodigal v2.6 (Hyatt et al. 2010). Clustering of the predicted coding sequence (CDS) was performed with CD-HIT-EST using the settings “-C 0.95, -T 0.8” (Li & Godzik 2006). Identification of unique and shared clusters were done using basic unix commands e.g. csplit, grep, sort and uniq. The specific commands used and files generated during clustering can be found in the Zenodo database (https://doi.org/10.5281/zenodo.1489356).

Phylogenetic analysis
Reconstruction of the Agrobacterium phylogeny used PhyloPhlAN (Segata et al. 2013). PhyloPhlAN is a bioinformatic pipeline that identifies conserved proteins (400 markers) from microbial genomes and uses them to construct a high-resolution phylogeny using maximum likelihood inference approach (Price et al. 2010). For single gene tree construction, protein sequences were aligned with mafft v7.3 (Katoh & Standley 2013) using the the most accurate setting (--localpair --maxiterate 1000) followed by phylogenetic tree construction via IqTree v1.65 with optimized model (Kalyaanamoorthy et al. 2017; Nguyen et al. 2014). Visualization and annotation of phylogenetic trees was performed with Figtree v1.4.3 (http://tree.bio.ed.ac.uk/software/figtree/).

Pan-genome construction and phylogenomics
Whole genome sequences were reannotated with Prokka v1.1 using the default setting (Seemann 2014). The Prokka-generated gff files were used as the input for Roary v3.12.0 to calculate the pan-genome (Page et al. 2015). Maximum likelihood tree construction of the core-genome alignment and tree visualization used FastTree2 v2.1.10 (-nt -gtr) (Price et al. 2010) and FigTree v 1.4.3, respectively. Input and output files associated with the Roary analysis have been deposited in the Zenodo database (https://doi.org/10.5281/zenodo.1489356).

Detection and visualization of Ti plasmid
Genome sequences of each member of the genomospecies 4 except for the problematic DSM 37014\textsuperscript{T} strain were used as the query for blastN search (-evalue 1e-100) against the octopine-type Ti plasmid (Altschul et al. 1990). The result of the similarity search was subsequently visualized in Blast Ring Image Generator (BRIG) v0.95 (Alikhan et al. 2011).

**Genome annotation and KEGG pathway reconstruction**

Whole genome sequences of *A. tumefaciens* B6\textsuperscript{T} and *A. radiobacter* NCPPB 3001\textsuperscript{T} were submitted to the online server GhostKoala (Kanehisa et al. 2016b) for annotation and the annotated genomes were subsequently used to reconstruct KEGG pathways (Kanehisa et al. 2016a) in the same webserver. Identification of proteins with TIGRFAM signatures of interest (Haft et al. 2003) used HMMsearch v3.1b2 with the option “--cut_tc” activated to filter for only protein hits passing the TIGRFAM trusted cutoff values (Johnson et al. 2010).

**RESULTS**

**An improved *Agrobacterium radiobacter* type strain genome**

Raw sequencing data and whole genome assembly for strains B6 and NCPPB3001 reported in this study are linked to the NCBI Bioproject IDs PRJNA300485 and PRNA300611, respectively. The newly assembled genome of *A. radiobacter* type strain that was sourced from the National Collection of Plant Pathogenic Bacteria (NCPPB) is approximately 30% smaller than the first reported *A. radiobacter* DSM 30147\textsuperscript{T} genome with 96% less contigs (22 vs 612), 20-fold longer N50 (480 kb vs 23 kb) and assembled length that is much more similar to other *Agrobacterium* spp. (Table 1). In addition, it is near-complete with 685 out of 686 BUSCO Rhizobiale single-copy genes detected as either partial or complete with minimal evidence of contamination as indicated by the near absence of duplicated single-copy gene(<0.1%). On the contrary, the current DSM 30147 genome is missing 25.1% of the single copy gene with up to 34.8% duplication rate. At the time of this manuscript writing, another genome of *A. radiobacter* type strain that was sourced from another culture collection centre e.g. the Belgian Coordinated Collections of Microorganisms has been deposited in the NCBI wgs database (*A. radiobacter* LMG140\textsuperscript{T}, Table 1) with assembly statistics that are highly similar to the type strain genome reported in this study.

**The inflated genome size of *Agrobacterium radiobacter* DSM 30147(T) is due to technical errors**

Instead of sharing a recent common ancestor as would be expected for a recently duplicated gene, the duplicated single copy genes coding for seryl-tRNA synthetase in *A. radiobacter* DSM 30147\textsuperscript{T} were placed in two distinct clusters with one affiliated to genomospecies 4 and the other affiliated to genomospecies 7 (Figure 1A). Such an unexpected clustering pattern raises the suspicion of genome assembly from two or more non-clonal bacterial strains. In addition, by performing comparison at the genome-scale based on whole proteome clustering of *A. radiobacter* DSM 30147\textsuperscript{T} /NCPPB
3001<sup>T</sup> (Previous study, GCF_000421945; This study, GCF_001541305), A. sp. TS43 (unpublished, GCF_001526605) and A. tumefaciens B6 (GCF_001541315), we observed a high number of proteins that were exclusively shared between Zhang et al A. radiobacter DSM 30147 and A. sp. TS43 belonging to genomospecies 7 (Figure 1B). Coincidentally, despite not sharing the same Bioproject ID, the whole genomes of strains DSM 30147<sup>T</sup> and TS43 were sequenced by the Zhang et al., and submitted to NCBI on the same date, 30-May-2013, hinting strain contamination during sample processing in the lab.

**Genome-scale average nucleotide identity calculation supports the amalgamation of A. radiobacter and A. tumefaciens into a single genomospecies**

Single gene tree shows that A. radiobacter NCPPB 3001<sup>T</sup> and A. tumefaciens B6<sup>T</sup> belong to the genomospecies 4 clade (Figure 1A), corroborating with the PhyloPhIaN phylogenomic tree that was constructed based on the alignment of 400 universal single-copy proteins (Figure S1). The pairwise average nucleotide (ANI) among strains within this clade is consistently more than 95% further supporting their affiliation to the same genomospecies (Figure 2) (Coutinho et al. 2016; Jain et al. 2017). As expected, pairwise ANI of less than 92% was observed when they were compared with strains from genomospecies 7 (strains RV3 and Zutra 3/1). A 100% pairwise ANI was observed between A. radiobacter type strains that were sourced from NCPPB and LMG. In addition, non-type strains B140/95 and CFBP5621 also exhibit a strikingly high pairwise ANI (>99%) to the type strains of A. tumefaciens and A. radiobacter, respectively, leading to the formation of sub-clusters within genomospecies 4 (Figure 2).

**Is A. radiobacter a non-tumorigenic strain of A. tumefaciens?**

A majority of the currently sequenced strains from genomospecies 4 are non-tumorigenic as evidenced by the near complete lack of genomic region with significant nucleotide similarity to the octopine-type Ti reference plasmid (Figure 3). Of the 14 genomes analyzed, only strains B6<sup>T</sup> and B140/95 exhibit a complete coverage of the Ti plasmid with near 100% sequence identity while strain 186 shows hits mainly to the essential gene clusters of a Ti plasmid such as the vir gene cluster (black rings and gene labels in Figure 3) at a substantially lower sequence identity (50%<x<90%) (Figure 3), suggesting that it may be harboring a dissimilar variant of Ti plasmid e.g. different opine type. In addition, although lacking hits to the virulence gene of the Ti plasmid, the tra and trb clusters involved in plasmid conjugal transfer are present in strains Kerr 14, CCNWGS0286 and UNC420CL41Cvi. Despite belonging to the same genomospecies, core genome alignment and phylogenomic analysis indicates that A. radiobacter NCPPB3001<sup>T</sup> is sufficiently divergent from A. tumefaciens B6<sup>T</sup> leading to their separation into two distinct sub-clusters (Figure 4A). This is also resonated by their different sub-cluster placement in the pairwise ANI heatplot (Figure 2). Furthermore, strains from both subclades could be broadly differentiated by the set of core accessory genes that they harbor (Figure 4B). Therefore, even though A. radiobacter does not harbor a Ti plasmid, it cannot be considered as a non-tumorigenic strain of A.
*tumefaciens* given multiple lines of evidence indicating its substantial genomic divergence from *A. tumefaciens*.

**Agrobacterium genomospecies 4 strains differ in their genomic potential for nucleotide sugar metabolism**

Individual comparison of the reconstructed KEGG pathways in *A. tumefaciens* (Figure 5A) and *A. radiobacter* (Figure 5B) revealed stark contrast in the anabolism of dTDP-L-rhamnose which is commonly found in the O-antigen of LPS in gram-negative bacteria. Surprisingly, the entire enzyme set required for the generation of dTDP-L-rhamnose from D-glucose-phosphate (Table 2) is absent in *A. tumefaciens* B6, suggesting that this common nucleotide sugar may be absent from the LPS O-antigen of strain B6. A manual inspection of the accessory genes uniquely shared by *A. tumefaciens* strains B6 and B140/95 identified a homolog cluster containing GDP-L-fucose synthase (EC 1.1.1.271) that is involved in the enzymatic production of GDP-L-fucose from GDP-4-dehydro-6-deoxy-D-mannose and NADH (Table 2 and Figure 5C). As expected, the genes coding for this enzyme and GDP-mannose 4,6-dehydratase involved in the conversion of GDP-alpha-D-mannose to GDP-4-dehydro-6-deoxy-D-mannose, are absent in the *A. radiobacter* NCPPB3001 genome (Figure 5D). Intriguingly, HMMsearch scan revealed the presence of two protein hits to the TIGR01479 HMM profile in *A. tumefaciens* B6 that corresponds to D-mannose 1,6-phosphomutase (EC 5.4.2.8) required for the synthesis of D-mannose 6-phosphate. In addition to strain B6, its close relative, strain B140/95, and a more distantly related strain Kerr14 also harbor two copies of this gene. However, one of the D-mannose 1,6-phosphomutases in strain Kerr14 is more divergent with a lower TIGRFAM HMM sequence score (Table 2). Furthermore, it exhibits less than 70% protein identity to the *A. tumefaciens* B6 and B140/95 homologs, forming a private protein cluster in the pan-genome (data not shown).

**DISCUSSION**

We re-sequenced the genome of *Agrobacterium radiobacter* type strain using strain directly obtained from NCPPB. The assembled *A. radiobacter* genome reported in this study exhibits assembly statistics that are consistent with a high-quality draft genome such as high genome completeness and contiguity, near-zero contamination/duplication and comparable genome size to other closely related strains (Gan et al. 2018; Parks et al. 2015). Furthermore, given the improved contiguity and dramatic reduction in the number of contigs of this newly assembled draft genome, we recommend using this genome in place of the previously published draft genome for future *Agrobacterium* comparative studies.

The distinct separation of *Agrobacterium* genomospecies 4 and 7 at 95% ANI cutoff corroborates with the previously established “genomic yardstick” for species differentiation (Konstantinidis & Tiedje 2005; Richter & Rosselló-Móra 2009). Using this percentage cutoff, the ANI approach has been successfully used to provide a near
“black-and-white” pattern of species separation in even some of the most diverse bacterial genera such as *Pseudomonas*, *Arcobacter* and *Stenotrophomonas* (Pérez-Cataluña et al. 2018; Tran et al. 2017; Vinuesa et al. 2018). Given the increasing evidence highlighting the robustness and reliability of the ANI approach in species delineation, the pairwise ANI between *A. tumefaciens* and *A. radiobacter* type strains that is at least 2.5% higher than the 95% cutoff value is rigorous evidence that they belong to the same genomospecies, effectively serving as the final nail in the coffin for the decade-long debate on their taxonomic status. The amalgamation of *A. radiobacter* and *A. tumefaciens* into a single species have been repeatedly suggested in the past few years but was complicated by the special status of *A. tumefaciens* as the type species of the genus *Agrobacterium* despite the priority that *A. radiobacter* has over *A. tumefaciens* as it was isolated and described 3 years before *A. tumefaciens* (Young et al. 2001; Young et al. 2003). Despite sharing numerous morphological and biochemical features, differences in genomic features such as pairwise ANI, phylogenomic clustering and core accessory gene contents do exist among members in *Agrobacterium* genomospecies 4 that can facilitate the identification of genotypic and phenotypic variants to accurately delimit sub-species relationships in the future (Brenner et al. 2015; Jezbera et al. 2011; Meier-Kolthoff et al. 2014; Tan et al. 2013).

To date the LPS for both type strains have been determined (De Castro et al. 2004; De Castro et al. 2002). In stark contrast to *A. radiobacter*, the *A. tumefaciens* LPS consists of D-arabinose and L-fucose that have yet been reported to date in another members of the genus *Agrobacterium* (De Castro et al. 2002). The presence of the L configuration of fucose is considered to be rare even among plant pathogenic bacteria but may be associated with the ability of *A. tumefaciens* to colonize or bind to wounded plant cell (Lippincott et al. 1977; Whatley et al. 1976; Whatley & Spiess 1977). It has been previously shown that the LPS of *A. tumefaciens* but not *A. radiobacter* can bind to the plant cells thus providing protection against subsequent infection by pathogenic strains (Whatley et al. 1976). The presence and absence of nucleotide sugars in the O-chain constituent of LPS in both type strains corroborates with their observed genomic potential in the nucleotide sugar metabolism pathway thus underscoring the utility of comparative genomics in facilitating the prediction of microbial host range and ecological niche (Klosterman et al. 2011). For example, the absence of L-rhamnose and L-fucose in the LPS of *A. tumefaciens* B6 and *A. radiobacter* DSM30147, respectively, is consistent with the lack of genes coding for enzymes involved with the particular nucleotide sugar metabolism. Generation of *Agrobacterium tumefaciens* B6 LPS mutant via targeted gene deletion (Kaczmarczyk et al. 2012) or the classical but more laborious transposon mutagenesis approach followed by characterization of the LPS mutant host-range and phytopathogenicity will be instructive (Gan et al. 2011; Reuhs et al. 2005).

Our current genomic sampling indicates that the Ti plasmid appears to be restricted to the *A. tumefaciens* subclade. The maintenance of the Ti plasmid is metabolically taxing given its large size (Barker et al. 1983; Glick 1995). Even if the Ti plasmid was conjugally transfer for example, to *A. radiobacter*, the inability of *A. radiobacter* to colonize plant host as evidenced by its LPS incompatibility will not confer
an advantage to the new plasmid host in a natural environment (Thomashow et al. 1980). Furthermore, in the absence of high density AHL signals which is required to trigger Ti plasmid conjugation (Fuqua & Winans 1994; Pappas 2008; Zhang et al. 2002), the newly acquired Ti plasmid in *A. radiobacter* may be cured in its natural soil habitat after a few generations. Although the spontaneous transfer of the Ti plasmid from tumorigenic *A. tumefaciens* to *A. radiobacter* K84 has been reported previously, strain K84 was re-classified based on a recent core gene analysis to *Rhizobium rhizogenes* (Velázquez et al. 2010; Vicedo et al. 1996), reiterating the pervasive taxonomic inconsistency within the genus *Agrobacterium* that may have confounded previous biological interpretations (De Ley et al. 1966; Lindström et al. 1995; Young 2008). Given that a large majority of *Agrobacterium* genetics was performed during the pre-NGS era (Gan & Savka 2018), it remains unknown as to how many *A. tumefaciens* and *A. radiobacter* strains have been molecularly misclassified due to their high genomic relatedness.

The inability to accurately identify plasmid and chromosomal-derived contigs among the draft genomes means that some of the core accessory genes among tumorigenic strains may be plasmid-derived and should be treated with caution as the low-copy-number Ti-plasmid is prone to curing in the absence of AHL signals. Despite the value of complete genome assembly in enabling the accurate partitioning of plasmid and chromosomal genomic region (Arredondo-Alonso et al. 2017), the representation of complete *Agrobacterium* genomes in current database is still very low as a majority of the genomes were assembled from short Illumina reads that cannot effectively span repetitive region (Wibberg et al. 2011; Wood et al. 2001). Furthermore, most *Agrobacterium* strains harbor multiple large plasmids that further complicate short-read-only assembly graph (Kado et al. 1981; Lowe et al. 2009; Shao et al. 2018). Given the currently available genomic resources for *Agrobacterium*, defining subspecies within the *Agrobacterium* genomospecies 4 based on the identification of lineage-specific gene set (Moldovan & Gelfand 2018) will be challenging. However, we anticipate that the advent of high throughput long-read sequencing that can span large repetitive region in recent years is likely going to overcome this limitation allowing a more accurate depiction of microbial pangenome (Gan et al. 2012; Gan et al. 2017; Schmid et al. 2018a; Schmid et al. 2018b). Future hybrid genome assemblies (Illumina and Nanopore/PacBio reads) of members from genomospecies 4 with comprehensive metadata and reliable phenotypic information, will be instructive.

CONCLUSIONS

Despite belonging to the same genomospecies, *A. tumefaciens* and *A. radiobacter* are by no means clonal at the chromosomal level and instead demonstrate sufficient genomic characters that qualify their separation into two sub-species. In addition, the difference in the LPS profile among two type strains will have implications to host specificity leading to geographical separation. In the spirit of preserving the naming of both species but at the same time respecting the taxonomic jurisdiction for strain priority, we propose *A. tumefaciens* to be reclassified as *A. radiobacter subsp.*
tumefaciens and for A. radiobacter to retains its species status with the proposed name of A. radiobacter subsp. radiobacter.

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Figure 1

Phylogenetic and genomic evidence indicating contamination in the published *A. radiobacter* DSM 30147T genome.

(A) Maximum likelihood phylogenetic tree of seryl-tRNA synthetases from Agrobacterium genomospecies 4 and 7. Codes after the tildes are contigs containing the corresponding homologs. Node labels indicate ultra-fast bootstrap support value and branch length indicates number of substitutions per site. Duplicated homologs in the problematic *A. radiobacter* DSM 30147 genome were colored red. (B) Venn diagram of the core proteome of selected Agrobacterium strains from genomospecies 4. Numbers in the overlapping regions indicate the number of coding sequences (CDS) that shared by two or more groups at 95% nucleotide identity cutoff.
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Figure 2

A heatmap showing the hierarchical clustering of *Agrobacterium* strains based on genomic distance.

Values in boxes indicate pairwise average nucleotide identity. Horizontal colored bar below the heatmap indicate the genomospecies assigned to each genome (G7, genomospecies 7; G4, genomospecies 4). Boxed labels indicate genomes sequenced in this study.
Figure 3

Prevalence and sequence conservation of the octopine-type Ti plasmid among Agrobacterium genomospecies 4.

Each genome (labelled 1-15) is represented by a colored ring shaded based on nucleotide percentage similarity to the reference Ti plasmid (min. 50%; max. 100%). The outermost ring highlights the gene regions involved in tumorigenesis (vir, iaa and ipt) and plasmid conjugation (trb and tra). Asterisks indicate genomes sequenced in this study.
Figure 4

Genomic divergence among genospecies 4 strains.

(A) Unrooted maximum likelihood tree constructed based on the core genome alignment. Branch length and node labels indicate number of substitutions per site and FastTree2 SH-like support values, respectively. Putative subclades were colored blue, red and purple (B) Distribution of accessory (non-core) gene clusters among strains determined with Roary and plotted with the perl script roary2svg.pl (https://github.com/sanger-pathogens/Roary/blob/master/contrib/roary2svg/roary2svg.pl). A total of 7,906 accessory gene clusters were identified by Roary and the number of accessory genes presence in each genome are shown in the most right column. Vertical grey lines/bars along the plot indicate presence of accessory gene. Asterisks indicate genomes sequenced in this study.
A

Unclassified subcluster

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B

|                  | Number of Accessory Genes |
|------------------|---------------------------|
| CFBP_5621        | 1463                      |
| LMG140           | 1629                      |
| NCPPB_3001*      | 1634                      |
| B140_95          | 1874                      |
| B6               | 1881                      |
| B6*              | 1792                      |
| 224MFTsu31       | 870                       |
| 719_389          | 990                       |
| UNC420CL41Cvi    | 1032                      |
| CCNWGS0286       | 1285                      |
| LMG215           | 1507                      |
| 186              | 1790                      |
| LAD9             | 1932                      |
| Kerr_14          | 1957                      |
Figure 5

KEGG pathway of nucleotide sugar metabolism associated with *Agrobacterium* lipopolysaccharide synthesis.

(A) and (B), genomic potential of *A. tumefaciens* B6 and *A. radiobacter* DSM 30147, respectively, in the biosynthesis of dTDP-L-rhamnose. (C) and (D), genomic potential of *A. tumefaciens* B6 and *A. radiobacter* DSM 30147, respectively, in the biosynthesis of GDP-L-Fucose. Numbers in boxes indicate Enzyme Commission numbers. White and green boxes indicate absence and presence of the corresponding enzymes, respectively, based on GhostKoala annotation (Kanehisa et al. 2016).
Table 1 (on next page)

Genome statistics of publicly available *Agrobacterium* genomospecies 4 whole genome sequences
Table 1: Genome statistics of publicly available Agrobacterium genomospecies 4 whole genome sequences

| Assembly accession | Strain | Isolation Source | Country   | Size | GC%  | # Contig |
|--------------------|--------|------------------|-----------|------|------|---------|
| GCF_900045375      | B6     | Apple Gall (Iowa) | USA   | 5.8  | 59.07 | 4       |
| GCF_001541315*     | B6     | Apple Gall (Iowa) | USA   | 5.6  | 59.32 | 52      |
| GCF_001692245      | B140/95| Peach/Almond Rootstock | USA | 5.7  | 59.23 | 45      |
| GCF_002179795      | LMG 215| Humulus lupulus gall (USA) | USA | 5.4  | 59.48 | 33      |
| GCF_000233975      | CCNWGS0286 | R. pseudoacacia nodules | China | 5.2  | 59.53 | 49      |
| GCF_900011755      | Kerr 14= LMG 15 = CFBP 5761 | Soil around Prunus dulcis | Australia | 5.9  | 59.04 | 5       |
| GCF_002591665      | 186    | English Walnut gall | California | 5.7  | 59.42 | 22      |
| GCF_002008215      | LMG 140 = NCPPB 3001 =CFBP 5522= DSM 30147 | saprobiic soil | Germany | 5.5  | 59.34 | 22      |
| GCF_000421945      | LMG 140 = NCPPB 3001 =CFBP 5522= DSM 30147 | saprobiic soil | Germany | 7.17 | 59.86 | 612     |
| GCF_001541305*     | LMG 140 = NCPPB 3001 =CFBP 5522= DSM 30147 | saprobiic soil | Germany | 5.5  | 59.36 | 22      |
| GCF_900012605      | CFBP 5621 | Lotus corniculata, root tissue commensal | France | 5.4  | 59.32 | 3       |
| GCF_003031125      | LAD9 (CGMCC No. 2962) | landfill leachate treatment system | China | 5.9  | 59.13 | 49      |
| GCF_000384555      | 224MFTsu31 | rhizosphere of L. luteus in Hungary, formerly R. lupini H13-3 | USA | 4.8  | 59.73 | 21      |
| GCF_900188475      | 719_389 | Rhizosphere and endosphere of Arabidopsis thaliana. | USA | 4.9  | 59.73 | 18      |
|   | GCF_000384555 | UNC420CL41Cvi | Plant associated | USA    | 5 | 59.69 | 18 |
|---|---------------|---------------|------------------|--------|---|-------|----|
| 1 | *Reported in this study |               |                  |        |   |       |    |
Table 2 (on next page)

Identification of *Agrobacterium* proteins with TIGRFAM domains involved in the biosynthesis of nucleotide sugar.

Numbers indicate bit scores calculated based on protein alignment to the model with higher scores indicating stronger and more significant hits.
Table 2. Identification of *Agrobacterium* proteins with TIGRFAM domains involved in the biosynthesis of nucleotide sugar. Numbers indicate bit scores calculated based on protein alignment to the model with higher scores indicating stronger and more significant hits.

| Assembly ID | Strain    | TIGR01479 (EC 5.4.2.8) | TIGR01472 (EC 4.2.1.47) | TIGR01207 (EC 2.7.7.24) | TIGR0118 (EC 4.2.1.46) | TIGR0122 (EC 1.1.1.133) | TIGR0121 (EC 5.1.3.13) | 1st hit | 2nd hit |
|-------------|-----------|------------------------|-------------------------|-------------------------|------------------------|------------------------|------------------------|---------|---------|
| GCF_9000453 | B6        | 690.2                  | 566.6                   | 589.5                   |                       |                        |                        |         |         |
| GCF_0015413 | B6        | 690.2                  | 566.6                   | 589.5                   |                       |                        |                        |         |         |
| GCF_0016922 | B140/95   | 690.2                  | 566.6                   | 589.5                   |                       |                        |                        |         |         |
| GCF_9000117 | Kerr14    | 691.3                  | 690.2                   | 428.6*                  |                       |                        |                        |         |         |
| GCF_0015413 | NCPPB3001 | 690.2                  | 494.6                   | 488.5                   | 215.4                 | 331.5                 |                        |         |         |
| GCF_0020082 | LMG140    | 690.2                  | 494.6                   | 488.5                   | 215.4                 | 331.5                 |                        |         |         |
| GCF_9000126 | CFBP5621  | 689.3                  | 494.6                   | 489.5                   | 215.4                 | 331.5                 |                        |         |         |
| GCF_0025916 | 186       | 689.3                  | 494.6                   | 488.5                   | 215.4                 | 331.8                 |                        |         |         |
| GCF_0030311 | LAD9      | 688.5                  | 494.4                   | 487.9                   | 215.4                 | 329.9                 |                        |         |         |
| GCF_002339  | CCNWGS    | 644.8                  | 494.6                   | 487.5                   | 215.4                 | 331.8                 |                        |         |         |
| GCF_0021797 | LMG215    | 690.2                  |                        |                        |                       |                       |                        |         |         |
| GCF_0003845 | 224MFTsu31| 644.8                  |                        |                        |                       |                       |                        |         |         |
| GCF_0004822 | UNC420CL41| 644.8                  |                        |                        |                       |                       |                        |         |         |
| GCF_9001884 | 719_389   | 687.5                  |                        |                        |                       |                       |                        |         |         |

*Formed a separate protein cluster from the rest of genomospecies 4 GDP-mannose-4,6-dehydratase orthologs (<70% pairwise protein identity)