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
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email: is@aber.ac.uk



Complete genome sequence of the biocontrol agent *Serratia marcescens* strain N4–5 uncovers an assembly artefact

Larissa Carvalho Ferreira^{1,2} · Jude E. Maul³ · Marcus Vinicius Canário Viana⁴ · Thiago Jesus de Sousa⁴ · Vasco Ariston de Carvalho Azevedo⁴ · Daniel P. Roberts³ · Jorge Teodoro de Souza² 

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Abstract

Serratia marcescens are gram-negative bacteria found in several environmental niches, including the plant rhizosphere and patients in hospitals. Here, we present the genome of *Serratia marcescens* strain N4–5 (=NRRL B-65519), which has a size of 5,074,473 bp (664-fold coverage) and contains 4840 protein coding genes, 21 RNA genes, and an average G + C content of 59.7%. N4–5 harbours a plasmid of 11,089 bp and 43.5% G + C content that encodes six unique CDS repeated 2.5× times totalling 13 CDS. Our genome assembly and manual curation uncovered the insertion of two extra copies of the 5S rRNA gene in the assembled sequence, which was confirmed by PCR and Sanger sequencing to be a misassembly. This artefact was subsequently removed from the final assembly. The occurrence of extra copies of the 5S rRNA gene was also observed in most complete genomes of *Serratia* spp. deposited in public databases in our comparative analysis. These elements, which also occur naturally, can easily be confused with true genetic variation. Efforts to discover and correct assembly artefacts should be made in order to generate genome sequences that represent the biological truth underlying the studied organism. We present the genome of N4–5 and discuss genes potentially involved in biological control activity against plant pathogens and also the possible mechanisms responsible for the artefact we observed in our initial assembly. This report raises awareness about the extra copies of the 5S rRNA gene in sequenced bacterial genomes as they may represent misassemblies and therefore should be verified experimentally.

Keywords *Serratia marcescens* · Artefacts · 5S rRNA · Biological control · Complete genome · Plant-associated bacteria

Data deposition: Trimmed sequence data and assembly are deposited in GenBank (accession numbers: NZ_CP031316.1 and NZ_CP031315.1).

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✉ Jorge Teodoro de Souza
jorge.souza@ufla.br

¹ Plant Pathology Department, Federal University of Lavras, Lavras, MG 37200-000, Brazil

² Institute of Biological, Environmental and Rural Sciences, Aberystwyth University, Aberystwyth SY23 3DA, UK

³ Sustainable Agricultural Systems Laboratory, USDA-Agricultural Research Service, Beltsville, MD 20705, USA

⁴ Institute of Biological Sciences, Federal University of Minas Gerais, Belo Horizonte, MG 31270-901, Brazil

Introduction

Soil-borne plant pathogens cause diseases that result in major reductions in crop yields [1]. These diseases are typically controlled in conventional crop production systems with strategies that include chemical pesticides [2]. Biologically based methods, such as the use of microbial biological control agents, are being developed to control these soil-borne pathogens due to problems associated with the availability and effectiveness of chemical pesticides and concerns regarding the impact of these chemicals on the environment and human health [3]. Microbes control plant diseases via several mechanisms, including predation, where the biological control agent produces an assortment of enzymes such as chitinases, proteases, and glucanases that degrade pathogen cell wall and other cellular components [4, 5]. Biological control agents can also produce antibiotics and other inhibitory molecules that kill or slow growth of the pathogen and can compete with the pathogen for resources such as nutrients and space.

Finally, certain biological control agents have been shown to associate with plants and induce defence responses that protect the plant from diseases [1].

The bacterium *Serratia marcescens* is ubiquitous in the environment and has been detected in association with plants [6] and animals [7] including humans in hospital settings [8], soil [9], water [10], and air [11]. Live cells and cell-free extracts of *S. marcescens* strains isolated from the environment have been shown to be effective in controlling certain soil-borne plant pathogens [12–14].

Here, we present the genome of *S. marcescens* N4–5, a strain isolated from soil and studied since 1996 [12] due to its effectiveness against multiple plant pathogens, including *Magnaporthe poae*, *Pythium ultimum*, and *Rhizoctonia solani* and its antimicrobial properties [12–15]. Strain N4–5 and its natural products, applied as seed treatments in biocontrol strategies, control seed, and seedling disease of cucurbits caused by the soil-borne plant pathogen *P. ultimum* [13, 15]. Among all the *S. marcescens* complete genomes available, only a few are from plant beneficial strains. Therefore, the addition of the N4–5 complete genome sequence into public databases will allow comparative analysis to better understand the mechanisms by which *S. marcescens* associates with plants and controls plant diseases, as well as the variety of lifestyles presented by this genus. Furthermore, we uncovered an assembly artefact in the genome of strain N4–5.

Material and methods

Genome sequencing and assembly

Strain N4–5 was obtained from New Jersey (USA) soil samples [12]. Genomic DNA was extracted with the QIAGEN Blood & Tissue genomic DNA isolation kit, using the manufacturer's protocol. The indexed library was constructed using Nextera® XT Index Kit v2 Set A, and the sequence data was generated in an Illumina NextSeq-500 using the run kit Illumina NextSeq® 500/550 High Output Kit v2. Sequencing resulted in 22,789,104 reads, with length varying from 32 to 151 bases, with an average length of 148 bp, where 94% of the reads contained 148–151 bases. The sequenced reads comprised a total of 3,369,822,757 bases and represented 664-fold genome coverage. The quality was checked with the programme FastQC v0.11.5 [16]. The paired-end library genome (2 × 149) was assembled using the SPAdes assembler available in PATRIC (Pathosystems Resource Integration Center) [17]. The 1634 contigs generated were united into 19 scaffolds using the CONTIGuator [18] with *S. marcescens* strain B3R3 (accession number CP013046.2) as the reference genome. Finally, gaps were closed with FGAP [19], NCBI's BLASTn [20], and read mapping in CLC Genomics Workbench 7.

The plasmid was assembled with reads that did not map against the final N4–5 complete nucleotide sequence using the programme plasmidSPAdes. The plasmid contigs were scaffolded using the *S. marcescens* strain A4Y201 plasmid pG5A4Y201 (accession number KJ541069.1) as reference. The plasmid sequence was finalized using the aforementioned FGAP, NCBI's BLASTn, and CLC Genomics Workbench. The structural and functional annotation was conducted as described above.

Genome annotation, manual curation, and analyses

The N4–5 genome was annotated using the RASTtk annotation service in PATRIC [21]. Manual curation was conducted through Artemis 16.0.0 software [22]. Translated protein sequences were confirmed with BLASTp against the UniProt database [23]. Clusters of Orthologous Groups (COGs) were inferred with the eggNOG v. 4.5.1 database [24]. Circular maps were generated using GCView Comparison Tool [25].

Manual curation revealed three copies of the 5S rRNA gene. To determine the veracity of this feature, PCR using primers designed on the regions flanking the 5S genes, MetA1F (5'- ACC GCA GGT AAC TCA TCA GG -3') and 23S1R (5'- GAC GTT GAT AGG CTG GGT GT- 3'), followed by sequencing with the Sanger method were performed as previously described [15]. The 100 bp DNA ladder (New England BioLabs) was used to visualize the band in the gel.

Phylogenomics and chemotaxonomic classification

Digital DNA-DNA Hybridization (dDDH) and Average Nucleotide/Amino Acid Identity (ANI/AAI) comparisons were calculated using GGDC [26], JspeciesWS [27], and Kostas Lab [28].

For FAME analyses, isolate N4–5 was grown for 24 h at 28 °C on trypticase soy broth agar (TSBA), and the composition of cellular fatty acid was determined by gas chromatography. Extraction and analyses were performed according to the recommendations of the MIDI (Microbial Identification) system.

Results and discussion

Genomic features

The *S. marcescens* N4–5 genome comprised a single chromosome of 5,074,473 bp, with 59.7% G + C content and a naturally occurring plasmid (Fig. 1). The chromosome had 4884 protein-coding genes, of which 4020 genes were functionally assigned, while the remaining genes were annotated as hypothetical proteins (Table 1). The N4–5 genomic nucleotide sequence contained 2747 transcription units and 992 operons.

Table 1 Genome features of *S. marcescens* strain N4–5

Attribute	Value
Chromosome size (bp)	5,074,473
N50	549,421
L50	4
GC (%)	59.69
Chromosomal genes	4884
Protein coding genes	4840
Plasmid (bp)	11,089
Genes in the plasmid	13
RNA genes	103
Pseudo genes	44
Genes with function prediction	4020
CRISPR repeats	2

From the 4884 genes, 3604 (73.8%) were classified in 22 functional COG categories. The most numerous COGs contained genes with general prediction only (408 genes), no function prediction (393 genes), and 401 genes involved in amino acid transport and metabolism, whereas the COG categories with the least number of genes contained one gene for RNA processing and modification and one for chromatin function and dynamics (Fig. 1b).

The circular plasmid comprised 11,089 bp and had 43.5% G + C content. The size of the plasmid was confirmed by digestion with restriction enzymes followed by electrophoresis. The plasmid sequence encoded six unique CDS that were repeated 2.5× totalling 13 CDS. From the six unique CDS, four were annotated as hypothetical proteins (Fig. 1c).

Biocontrol and plant-beneficial traits

Strain N4–5 is a known producer of the broad-spectrum antimicrobial prodigiosin, which contributes to its biological control activity [13, 15]. In accordance, the genome of N4–5 harboured the 14 canonical genes for prodigiosin biosynthesis (pig cluster, pigA–N) described by [29, 30]. As seen in other bacteria [30], the N4–5 pig cluster was flanked by *copA* and *cueR* homologues; however, differently from the other studied strains, N4–5 has a putative membrane protein (41 amino acids) annotated between *pigA* and *cueR*. Sixty-nine multidrug resistance genes were found during functional annotation of the N4–5 genome, including resistance to kasugamycin, biocyclomycin, fosmidomycin, and fusaric acid, which are antibiotics produced by microorganisms. Strain N4–5 also harbours chitinase genes (*chiA*, *chiB*, *chiD*, and *chiA1*) in the genome.

Genome analysis revealed that N4–5 encodes the siderophore enterobactin gene cluster containing *entA*, *entB*, *entC*, *entE*, *entF*, and *entH*, but the vibriobactin genes were absent. Furthermore, N4–5 carried 16 *tonB*-dependent

transporter genes, which are cellular receptors of siderophores. The production of siderophore complexes by bacteria contributes to enhance plant growth as they sequester iron from the environment and make it available for plant uptake [31]. The ability to utilize carbon sources provides a fitness advantage during microbial competition. The N4–5 genome had 267 genes responsible for carbohydrate transport and metabolism, comparable with *Pseudomonas alcaliphila* JAB1, a degrader of organic pollutants that had 196 genes with this functionality [32]. The surfactant serrawettin W1 was coded by one NRPS (non-ribosomal peptide synthase) gene with 3936 bp. Serrawettin W1 has antimicrobial, antitumor, and zoosporicidal activities and has potential uses in agriculture, medicine, and industry [33]. Altogether, these genome features support N4–5 as a potential biocontrol agent as well as a plant-beneficial strain.

Phylogenomics and chemotaxonomic data

Phylogenetic trees constructed with whole genome sequences and with sequences of the 16S gene placed strain N4–5 within the *S. marcescens* clade (Supplementary Fig. S1 and S2). The identity of these seven copies of the 16S rRNA gene found in the genome of strain N4–5 varied from 99.7 to 100%, indicating that strain N4–5 possesses low intragenomic variation in the ribosomal genes. Further analyses, including Average Nucleotide/Amino Acid Identity (ANI/AAI) and digital DNA-DNA Hybridization (dDDH) confirmed the classification of strain N4–5 as *S. marcescens*. The values for ANI, AAI, and dDDH were above the cut off for species delineation, 95, 95, and 70%, respectively, when compared with other *Serratia* genomes (Table S1).

The major components of the *S. marcescens* N4–5 fatty acid profile were C16:0 (22.4%), C17:0 cyclo (12.13%), C10:0 3OH (12.07%), and C12:0 3OH (5.15%). Minor fatty acid components were identified at less than 5%, most of which being common among previously identified species within the genus *Serratia*: C12:0, C12:0 2OH, C14:0, C14:0, C18:0, and C19:0 cyclo w8c. Some fatty acids isolated from N4–5 co-occurred in just a few other species. For example, C10:0 3OH was only shared with *S. plymuthica* and *S. rubidaea*, whereas C12:1 3OH, C12:0 3OH, C14:0 2OH, and C17:0 were only identified in other *S. marcescens*-GC subgroups.

Genome assembly artefact

Two extra copies of the rRNA 5S gene were found in the fifth ribosomal cluster of strain N4–5 after the initial assembly performed in SPAdes. These extra copies of the 5S rRNA gene were also present in all the other four assemblies we performed with data from different sequencing lanes with coverage 160–167×. The combined assembly, with 664× coverage was chosen due to the best metrics it returned, namely,

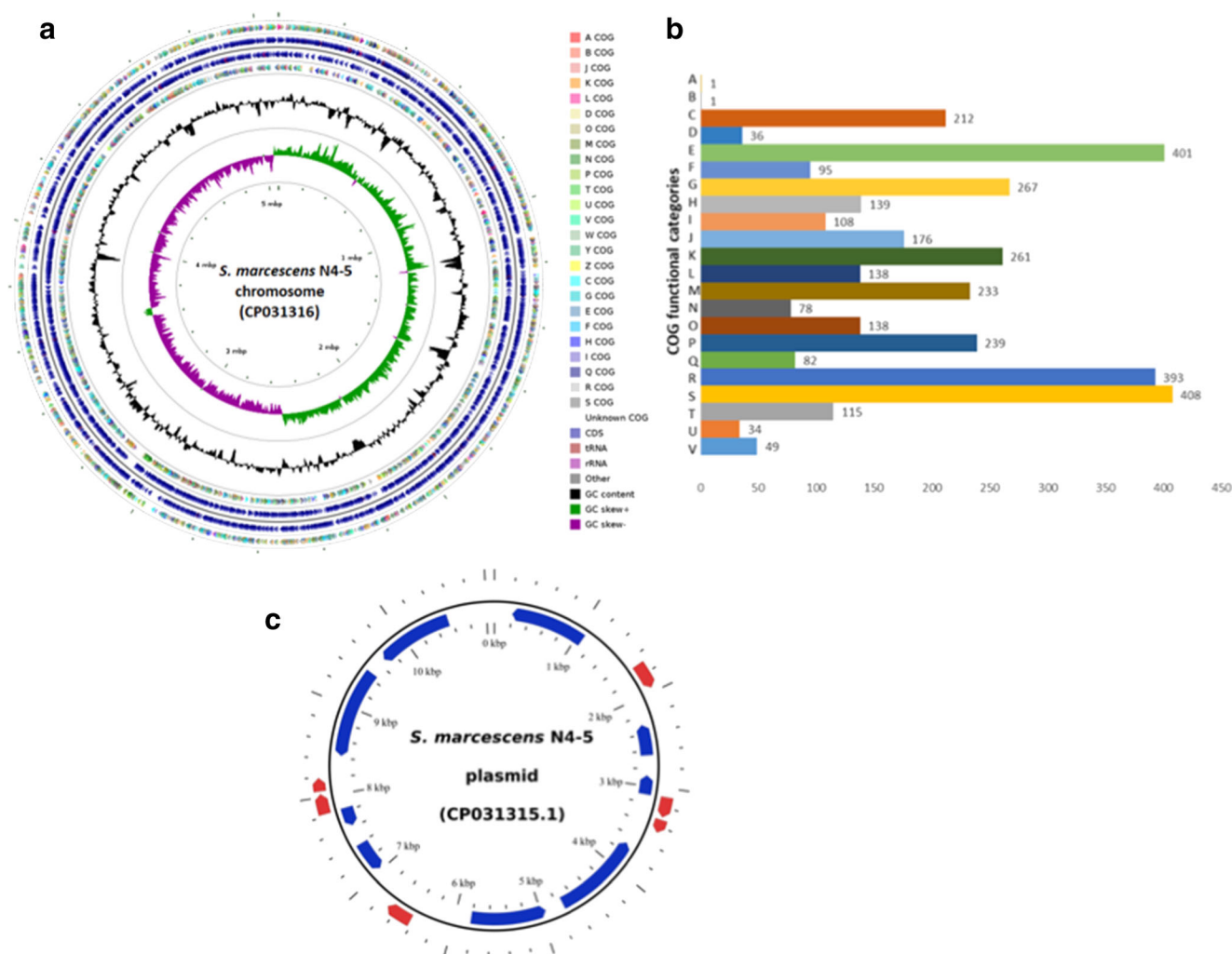


Fig. 1 *Serratia marcescens* strain N4-5 genome features. **(a)** Chromosome of N4-5 showing from outer circle to the centre: CDS on forward strand (coloured according to COG categories), all CDS and RNA genes on forward strand, all CDS and RNA genes on reverse strand,

CDS on reverse strand (coloured according to COG categories), G + C content and GC skew. **(b)** COG functional classification of strain N4-5 proteome. **(c)** Circular map of N4-5 plasmid showing the CDSs

highest N50 and N75, lowest L50 and L75, largest contig size, and overall total length assembled. To verify whether it was an artefact or a natural feature in strain N4-5, PCR amplification and sequencing with the Sanger method of the rRNA 5S gene region were performed. The primers were anchored in the gene *metaA* and in the 23S rRNA gene (Fig. 2). The 755 bp sequence obtained was mapped to the 989 bp region in the assembly and unequivocally showed that these extra copies of the 5S rRNA gene were assembly artefacts. The genome sequence was corrected accordingly, and therefore, strain N4-5 had regular ribosomal operons, i.e. one copy of the 5S, 16S, and 23S genes per cluster (Fig. 2). In strain N4-5, the assembly process was responsible for generating the artefact, but we currently do not know how the programme produces them. One possibility, although we did not test this hypothesis, is the fact that we used a genome with an extra copy of the 5S rRNA gene as the reference.

Extra copies of the 5S gene were also found in 72 completely sequenced genomes of *Serratia* strains deposited in public databases (Supplementary Table S2). The extra copies of the 5S gene were found in the complete genomes of species in other genera, including *Serratia*, *Proteus*, *Vibrio*, *Yersinia* (Supplementary Table S3), and possibly other bacterial genera. There is limited information on the biological implications of extra copies of the 5S gene, and it is certainly a subject that deserves further investigation. Many of the genomes that contain extra copies of the 5S gene were sequenced by long reads technologies such as PacBio (Supplementary Table 2), which is supposed to obtain reads that encompass the whole ribosomal region. Nevertheless, the occurrence of the extra copies of the 5S rRNA gene in these genomes should be verified with sequencing technologies with read lengths longer than 600 bp.

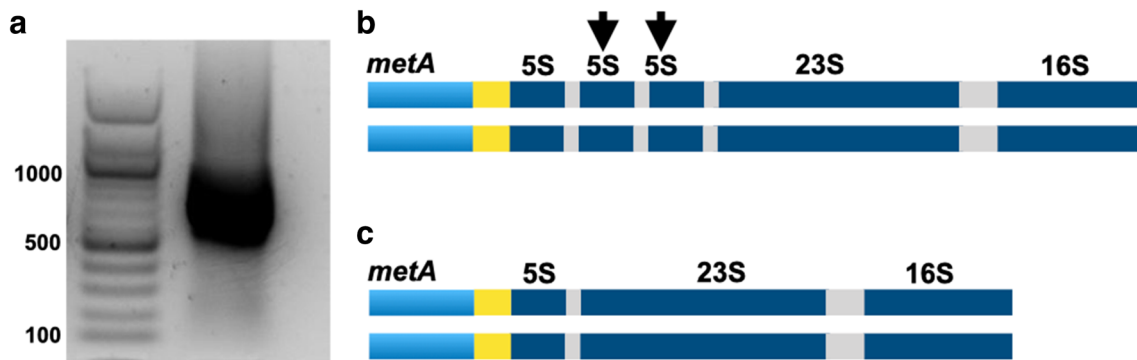


Fig. 2 Assembly artefact in the genome of *S. marcescens* strain N4-5. (a) Gel image representing the expected band size of 755 bp for a single 5S gene and its flanking regions. Band sizes are given in bp. (b) Schematic representation of the ribosomal operon with the assembly artefact in the genome of strain N4-5. Both DNA strands are represented. The rRNA genes are represented in dark blue, the gene *metA* is shown in light blue,

the spacers in the ribosomal operon in grey, and the intergenic region between *metA* and the ribosomal operon in yellow. The black arrows indicate the extra copies of the 5S gene in both DNA strands. This genomic region is not drawn to scale. (c) Ribosomal operon without the assembly artefact in strain N4-5

The number of copies of the 5S gene in bacterial genomes is thought to be identical to the number of copies of the other genes (23S and 16S) in the ribosomal operon. This assumption is supported by the fact that ribosomal genes and multigene families are homogenized by recombination through concerted evolution [34]. However, an estimate by rmDB, a database that documents the number of rRNA and tRNA genes in bacteria and archaea, revealed that 23.6% of the bacterial genomes have unequal copies of the rRNA genes, due mainly to additional copies of the 5S rRNA gene [35]. In the genus *Serratia*, this unequal number of 5S rRNA genes occurs in 96% of the sequenced genomes (Supplementary Table 2). It would be interesting to verify if these numbers are real or are artefacts caused by the in silico assembly programmes. The discovery and correction of assembly errors in draft genomes is a crucial problem that persists in Eulerian assemblies and genome assemblies in general [36].

Strain N4-5 is among the three *S. marcescens* with complete genomes in public databases without extra copies of the 5S rRNA gene. This does not mean that genomes without extra copies of the 5S rRNA genes are correctly assembled and the other ones are misassemblies. With these results, we want to emphasize the need to verify these in silico assemblies with standard laboratory experimental procedures.

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