



Draft genome sequence of an endophytic bacterium, *Paenibacillus tyrfis* strain SUK123, isolated from *Santiria apiculata* stem



Emmanuel Haruna^{a,b}, Noraziah M. Zin^{a,*}, Jonathan M. Adams^{c,*}

^a Programme of Biomedical Science, School of Diagnostic and Applied Health Sciences, Faculty of Health Sciences, Universiti Kebangsaan Malaysia, Kuala Lumpur, Malaysia

^b Department of Biochemistry Kaduna State University, Kaduna, Nigeria

^c Division of Agrifood and Environment, School of Water, Energy and Environment, Building 52a, Cranfield University, Cranfield, Bedfordshire MK43 0AL, UK

A B S T R A C T

Here we report the draft genome sequence of an endophytic *Paenibacillus tyrfis* strain isolated from the Universiti Kebangsaan Malaysia reserve forest, Malaysia. The genome size was approximately 8.04 Mb, and the assembly consisted of 107 scaffolds with 168 contigs, and had a G + C content of 53%. Phylogenetic analysis of strain SUK123 using the 16S rRNA gene revealed that it belonged to the family *Paenibacillaceae* with the highest similarity to *Paenibacillus elgii* SD^T (99%). Whole genome comparison of SUK123 with related species using average nucleotide identity (ANI) analysis revealed a similarity of 98% to *Paenibacillus tyrfis* Mst1^T, 94% to *Paenibacillus elgii* B69^T, 91% to *Paenibacillus ehimensis* A2^T, 68% to *Paenibacillus polymyxa* SC2^T and 69% to *Paenibacillus alvei* DMS29^T. The draft genome was deposited at the European Nucleotide Archive (PRJEB21373).

Specifications

Organism/cell line/tissue	<i>Paenibacillus tyrfis</i>
Sex	Not applicable
Sequencer or array type	Illumina Miseq
Data format	Raw data and analyzed i.e. assembled
Experimental factors	<i>Paenibacillus tyrfis</i> strain was isolated from <i>Santiria apiculata</i> stem
Experimental features	Isolation of bacteria, Genome sequencing, de novo assembly
Consent	Not applicable
Sample source location	Universiti Kebangsaan Malaysia reserve forest, Latitude & Longitude – 2.9125 & 101.7872

1. Direct link to deposited data

<http://www.ebi.ac.uk/ena/data/view/PRJEB21373>.

2. Introduction

The bacterial genus *Paenibacillus* has been isolated from many environments, mostly from plant organs and their surrounding soil [1]. The species of *Paenibacillus* was included in the genus *Bacillus* until 1993 when it was proposed that the member of the “16S rRNA group 3” bacilli be distinguished from it [2]. Presently, the genus *Paenibacillus* consists of 395 known species [3]. Most members of the genus *Paenibacillus* are gram variable, either aerobic or facultatively anaerobic, rod-shaped, and endospore-forming with peritrichous flagella for motility [4]. The DNA G + C content of the various species of *Paenibacillus* ranges between 39 and 54 mol% whilst anteiso-C_{15:0} is the major cellular fatty acid and meso-diaminopimelic acid is the cell wall peptidoglycan diamino acid [5]. Most members of this genus have been reported to be producers of either active antimicrobial metabolites such as lipopeptides [6,7], plant-growth promoting hormones [8,9], or insecticides [10].

3. Experimental design, materials and methods

The endophytic *Paenibacillus tyrfis*, strain SUK123, was isolated from

* Corresponding authors.

E-mail addresses: noraziah.zin@ukm.edu.my (N.M. Zin), j.m.adams@cranfield.ac.uk (J.M. Adams).

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Table 1
Statistics of assembled sequence length.

No. of all scaffolds	107
Bases in all scaffolds	8,041,385
No. of large scaffolds (> 1000 bp)	73
Bases in large scaffolds	8,024,609
Largest length	505,699
Scaffold N50	257,089
Scaffold N90	62,404
G + C content (%)	53.061
No. of all contigs	168
Bases in all contigs	8,041,324
No. of large contigs (> 1000 bp)	114
Bases in large contigs	8,013,898
Largest length	501,150
Contig N50	197,909
Contig N90	48,401

the *Santiria apiculata* stem located at the Universiti Kebangsaan Malaysia forest reserve, whilst screening for endophytes with antimicrobial potential against ESKAPE (*Enterococcus faecium*, *Staphylococcus aureus*, *Klebsiella pneumoniae*, *Acinetobacter baumannii*, *Pseudomonas aeruginosa*, and *Enterobacter species*) pathogens. After sample collection, the stem surface was sterilized to avoid surface-bound epiphytes as described previously [11]. The sterilized stem was cut into a 2 cm section and plated on water agar plate medium from where SUK123 (an antibiotic producing bacterium) was isolated.

Endophytic *Paenibacillus tyrfis* DNA was extracted using an Ultraclean Microbial DNA Isolation Kit as described by the manufacturer (Mo Bio Laboratories, 2746 Loker Ave W # A, Carlsbad, CA 92010, USA). The bacterium was identified by sequencing its 16S rRNA gene. Sequence was performed on Illumina Miseq platform (Majorbio, China) by 2 × 400 bp paired-end libraries. The raw sequence quality was assessed using PRINSEQ lite version 0.20.4 and the genome was assembled using SOAPdenovo v2.04 with all parameters set by default [12] and GapCloser v1.12 was used to fill local inner gaps. The open reading frames (ORFs) were predicted using Glimmer 3.02 [13]. The

biological functions of these predicted ORFs were annotated using various databases.

4. Data description

The analyses of the assembled genome revealed a genome size of about 8,041,385 bp made of 107 scaffolds, 168 contigs, and with a G + C content of 53.06% and N50 contigs size of 197,909 (Table 1). The genome was annotated using Rapid Annotation Subsystems Technology server [14].

The RAST server prediction revealed 7368 coding sequences (CDS) with a total of 2880 CDS (40%) constituting 2747 and 133 of non-hypothetical and hypothetical proteins respectively in the subsystem coverage. A total of 4488 CDS (60%) comprising 2003 and 2485 of non-hypothetical and hypothetical proteins respectively were outside the subsystem coverage (Fig. 1).

The average nucleotide identity (ANI) [15] analysis of SUK123 with closely related species, revealed a similarity index with *Paenibacillus tyrfis* Mst1^T (98.06%), *Paenibacillus elgii* B69^T (93.98%), *Paenibacillus ehimensis* A2^T (91.17%), *Paenibacillus alvei* DMS29^T (68.81%) and *Paenibacillus polymyxa* SC2^T (68.27%), suggesting our isolate most closely related to *Paenibacillus tyrfis* Mst1^T.

Declaration of interest

We declare no conflict of interest with respect to this article titled ‘Draft genome sequence of an endophytic *Paenibacillus tyrfis* strain SUK123 isolated from *Santiria apiculata* stem’ submitted to Genomics data journal.

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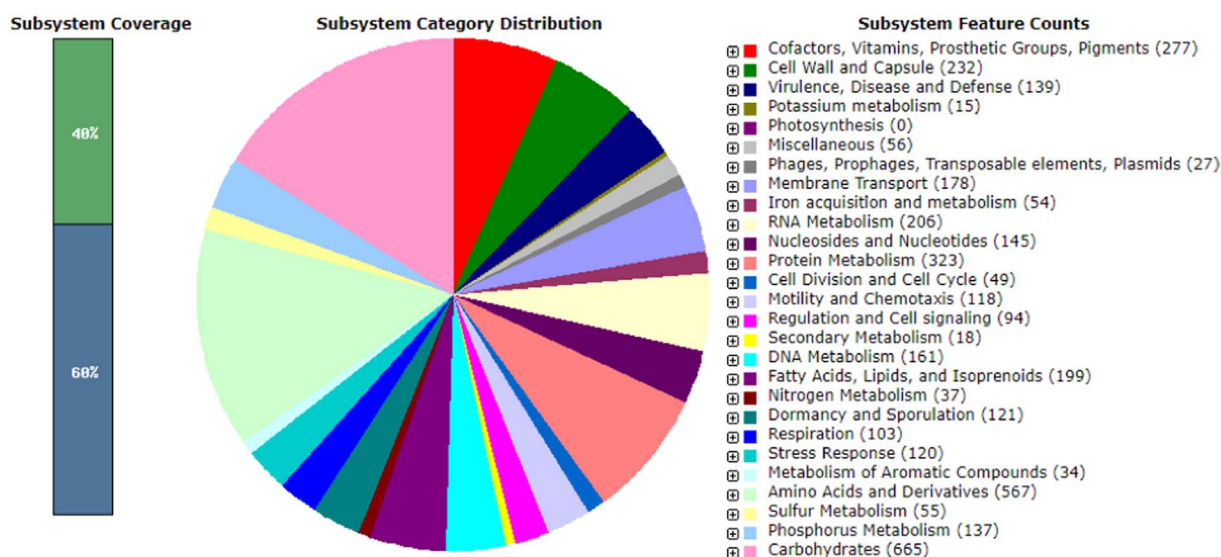


Fig. 1. A diagrammatic representation of *Paenibacillus tyrfis* strain genome subsystem coverage as annotated by RAST.

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