Whole Genome Sequencing of Sunflower Root-Associated *Bacillus cereus*

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**ABSTRACT:** In recent times, diverse agriculturally important endophytic bacteria colonizing plant endosphere have been identified. Harnessing the potential of *Bacillus* species from sunflower could reveal their biotechnological and agricultural importance. Here, we present genomic insights into *B. cereus* T4S isolated from sunflower sourced from Lichtenburg, South Africa. Genome analysis revealed a sequence read count of 7,255,762, a genome size of 5,945,881 bp, and G + C content of 34.8%. The genome contains various protein-coding genes involved in various metabolic pathways. The detection of genes involved in the metabolism of organic substrates and chemotaxis could enhance plant-microbe interactions in the synthesis of biological products with biotechnological and agricultural importance.

**KEYWORDS:** Bacterial endophytes, plant-microbe interactions, root architecture, secondary metabolites, siderophore

**INTRODUCTION**

Insights into plant endosphere communities have provided several opportunities in ensuring sustainable agriculture. Endosphere represents the entire internal tissue of plants colonized by microorganisms called ‘microbial endophytes’, which is beneficial to the host plants without eliciting any pathological effect. The role of microbial endophytes has been postulated with significant interest in agricultural biotechnology for plant growth and protection against environmental stress adaptors. Remarkably, identifiable bacterial species in the genus *Bacillus* from plant roots with plant growth-promoting characteristics and application in the greenhouse and field trials upon inoculation that enhances cucumber yield has suggested their potential in the formulation of bioinoculants in maintaining plant health and disease control.

Genetic modification of *Bacillus* species has contributed immensely to ecological balance with multifaceted significance in mitigating the excesses of chemical fertilizer application, and suppression of stress adaptors. The synthesis of bioinoculants from agriculturally important microbes has been employed as an alternative to develop environmentally friendly agriculture and to minimize threats posed by agro-chemical usage in food production. However, the rationale of bacterial dynamics and their mechanisms in plant growth-promotion has been explained, but the actual mechanisms are less understood.

The genomic analysis of *Bacillus* species associated with the medicinal plant, *Costus igneus* has been reported, nevertheless, genomic insights into *Bacillus* species associated with sunflower in Southern Africa has not been studied. Hence, studying the entire genomic structure of strain T4S from sunflower can pinpoint a delivery roadmap in the identification of novel genes participating in the metabolic pathways of biomolecules as the whole genome reports on *Bacillus* species in the endosphere of sunflower are rarely studied. Hence, integration of genomic data analysis will help predict important genes involved in metabolic pathways expressed by root-associated bacterial endophytes. Thus, this study presents the genomic analysis of endophytic bacterium *B. cereus* T4S earlier isolated from the sunflower root endosphere for improved sunflower yield.

The bacteria isolated from sunflower roots were sourced from farmlands in Lichtenburg, South Africa (26°4'31.266"S, 25°58'44.442"E) in February 2020. The healthy sunflower plants were carefully uprooted, placed inside sterile zip-lock bags, transported into the laboratory and stored at 4°C. The root samples were cut into small sizes with a sterile scalpel and washed in sterile distilled water. To ensure complete removal of the epiphytic bacteria, surface sterilization was achieved by soaking the samples in 70% ethanol for 3 minutes, followed by 3% sodium hypochlorite for 3 minutes, 70% ethanol for 30 seconds and rinsed with sterile distilled water. The level of sterility of the samples was assessed by pour plating on Luria-Bertani (LB) agar using the last water used to rinse plant samples. One gram (1 g) of plant material was weighed, suspended in 1 M phosphate–buffered solution (PBS) and manually macerated in a mortar and pestle until a smooth suspension was obtained. Sample suspensions were serially diluted up to 10^-9 dilutions and 0.1 mL of an aliquot from dilutions 10^-5 and 10^-6 were pipetted into Petri dishes and pour plated with sterilized LB agar. Inoculated Petri plates were incubated at 28°C for 24 hours. Distinct bacterial colonies formed on the plates were counted and selected based on morphological appearance. Pure culture of the bacterial isolate was obtained by repeated streaking onto sterile LB agar and incubated at 28°C for 24 hours. The pure bacterial strains were kept on an agar slant and stored at 4°C for further use.
The stored purified *B. cereus* T4S in 30% glycerol at ~80°C were sub-cultured on LB agar (g/L; NaCl – 10 g, tryptone – 10 g, and yeast extract – 5 g) and incubated at 28°C for 24 hours, then used for DNA extraction. Bacterial DNA was extracted by employing a commercial Quick-DNA™ Miniprep Kit specific for fungi or bacteria (Zymo Research, Irvine, CA, USA; Cat. No. D6005), following the manufacturer’s protocol. To quantify the purity of DNA, a NanoDrop spectrophotometer (Thermo Fischer Scientific, CA, USA) was used, while DNA quality was evaluated on 2% agarose gel and electrophoresed. The WGS of endophytic bacterial strain T4S was performed on Illumina’s Nextseq platform following the standard Illumina methods at Inqaba Biotechnical Industries (Pty) Ltd, Pretoria, South Africa.

Genomic sequences were analysed on the KBase platform.8 The quality assessment of the sequence reads was evaluated by FastQC (version 0.11.5),9 while the removal of sequence adapter and low-quality bases was processed with trimmomatic (version 0.36).10 Furthermore, sequence reads were assembled with SPAdes (version 3.13.0).11 Gene annotation and prediction were performed on RASTtk (Rapid Annotations using Subsystems Technology toolkit – version 1.073) and the publicly available NCBI (https://www.ncbi.nlm.nih.gov/) Prokaryotic Genome Annotation Pipeline (PGAP).12 All analyses were performed using default parameters unless otherwise specified. Secondary metabolites were determined by antiSMASH (version 6.0.0).13 The circular genome visualization was obtained from KBase (https://kbase.us/), while the phylogeny analysis was performed using MrBayes (http://www.phylogeny.fr/one_task.cgi?task_type=mrbayes) (version 3.2.6).8,14 The genomic comparison of strain T4S with other *B. cereus* was achieved as described by Zeng et al.15 In addition, in vitro screening of the plant growth-promoting potential of the strain T4S was performed using plate assay.

The strain T4S responded positively to almost all the plant growth-promoting tests (Table 1).

The comparison of genomic features of *B. cereus* strain T4S with other *B. cereus* is presented in Table 2. Notably, the whole-genome sequence analysis of strain T4S had a sequence read

### Table 1. Plant growth-promoting features of *B. cereus* T4S.

| PLANT GROWTH-PROMOTING TESTS | ISOLATE SCREENING | QUALITATIVE | QUANTITATIVE |
|-----------------------------|------------------|-------------|--------------|
| IAA                         | +                | 11.29 ± 0.01a |
| Siderophore                 | ++               | 87.30 ± 0.38c |
| Phosphate                   | +                | 30.43 ± 0.18b |
| Exopolysaccharide           | +++              | –            |
| Enzyme assay                | ZOC (mm)         |              |
| Amylase                     | –                | 0 ± 0.00a    |
| Cellulase                   | ++               | 50.00 ± 0.01d |
| Xylanase                    | ++               | 180.00 ± 0.01e |
| Mannanase                   | ++               | 35.00 ± 0.01c |
| Protease                    | +                | 4.00 ± 0.01b  |

Abbreviations: IAA, indole acetic acid; ZOC, zone of clearance measurement. Mean ± standard deviation values with different superscripts (small letters) within the same column represent a significant difference.

− = negative reaction.
+ = positive reaction.

### Table 2. Comparison of genomic features of *B. cereus* strain T4S with other *B. cereus*.

| GENOME | B. CEREUS T4S | B. CEREUS LCR12 | B. CEREUS 905 | B. CEREUS PG86311 | B. CEREUS AR156 | B. CEREUS UW85 |
|--------|---------------|-----------------|---------------|-------------------|-----------------|---------------|
| Genome size (bp) | 5 945 881 | 6 031 570 | 5 386 583 | 5 744 144 | 5 671 798 | 6 404 077 |
| G + C value | 34.8 | 34.8 | 35 | 35 | 35.25 | 34.81 |
| N₅₀ | 65 078 | 3706 | 91 | 5 230 609 | 5 160 326 | 285 |
| L₅₀ | 32 | 1 | 17 | 1 | 1 | 6 |
| Number of contigs | 198 | 40 | 127 | 3 | 4 | 54 |
| Number of genes | 6086 | 6078 | 5560 | 5801 | 5320 | 5321 |
| NCS | 6277 | 6005 | 5412 | 5625 | 5725 | 6257 |
| tRNA genes | 56 | 108 | 83 | 108 | 99 | 70 |
| rRNA genes | 5 | 48 | 6 | 42 | 39 | 38 |
| Bio-project numbers | SAMN18138757 | SAMN05436732 | SAMN04496851 | SAMN09783485 | SAMN04924832 | SAMN05171626 |
| Bio-sample | PRJNA706601 | PRJNA331062 | PRJNA312259 | PRJNA485192 | PRJNA320205 | PRJNA322638 |
| Accession number | JAFNAY000000000 | MCAX000000000 | LSTW000000000 | SAUQ000000000 | CP015589 | LYVD000000000 |

Abbreviation: NCS, number of the coding sequence.
count of 7,255,762, a genome size of 5,945,881 bp, and a G+C value of 34.8%. The average read length was 151 bp. The \( N_{50} \) and \( L_{50} \) values of the raw sequences were 65,078 and 32 bp, while the number of contigs and subsystems were 198 and 341, respectively (Table 2). Also, genome analysis revealed 6,277 coding sequences and 63 RNAs. Figure 1 shows the phylogeny of the sequence data of \( B. \) cereus T4S, while the subsystem category distribution of the key PCG is presented in Figure 2. The subsystem statistics showed 27 subsystem feature counts of the coding protein into functional groups with a total of 5159 PCG. The 1961 genes annotated by the SEED viewer were grouped into molecular function, cellular components and biological processes. The topmost 5 groups were protein metabolism (155 genes), vitamins, prosthetic groups, pigments (161 genes), cofactor, amino acids and derivatives (377 genes), carbohydrates (271 genes) and nucleosides and nucleotides (119 genes). The circular representation of the draft genome of \( B. \) cereus strain T4S is presented in Figure 3.

The secondary metabolite gene clusters, namely, non-ribosomal peptides (NRPS) and siderophore of strain T4S
Figure 3. Circular representation of the draft genome of *Bacillus cereus* strain T4S. Each color from the external to internal circle depicts green (GC skew +), and red (ORF). The ring black coloration (GC content) at the peak indicated higher or lower values than average GC content. The GC skew (−/+ ) in purple/green peaks in/outside the circle indicated values greater or smaller than 1. The GC skew is calculated as G-C/G+C.

Table 3. Information on the siderophore similar known gene cluster.

| SIDEROPHORE | Genes | Other | Other | Other | Additional | Core synthetic | Core synthetic |
|-------------|-------|-------|-------|-------|------------|----------------|----------------|
| Metabolites | Petrobactin biosynthesis (PB) | protein AsbE | PB aryl carrier protein AsbD | 3,4 dihydroxybenzoic acid-AMP ligase | AsbC | PB protein AsbB | PB protein AsbA |
| BA | 1986 | 1985 | 1984 | 1983 | 1982 | 1981 |
| Location | 7090-6248 | 6210-5227 | 5203-4928 | 4931-3693 | 3706-1868 | 1809-1 |
| BLAST hit with | J0658-00175 (30 471-31 320) | J0658-00180 (31 357-32 341) | J0658-00185 (32 364-32 631) | J0658-00190 (32 627-33 866) | J0658-00195 (33 852-35 691) | J0658-00200 (35 751-37 560) |
| Percentage identity | 88 | 93 | 93 | 93 | 84 | 92 |
| Percentage coverage | 100 | 100 | 103 | 100 | 100 | 100 |
| BLAST hit score | 506 | 631 | 158 | 784 | 1064 | 1052 |
| E-value | 2.4e-143 | 5.9e-181 | 5.1e-39 | 6.8e-227 | 6.6e-311 | 3.1e-307 |
| Function | Sugar phosphate isomerase/epimerase | Petrobactin biosynthesis protein AsbE | Petrobactin biosynthesis protein AsbD | AMP-building protein | Siderophore biosynthesis protein | luca/lucC family siderophore biosynthesis protein |
with 100% similarity selected and information on the functions of notable secondary metabolite (siderophore) genes (AsbABCDEF) that code for petrobactin synthesis/petrobactin-synthetic aryl carrier protein and 3,4 dihydroxybenzoic acid-AMP ligase, specific for additional biosynthetic, core synthetic and other genes were presented (Table 3).

Also, in Figure 4, the core structure of NRPS with a functional group, thiol (SH), carboxyl (COOH), amine (NH2), carbonyl (C=O) is represented. Genomic analysis revealed various genes involved in nitrogen fixation and nitrogen metabolism, iron transport and siderophore production, growth hormone synthesis, phosphate solubilization and transport, motility, biofilm formation and chemotaxis, biological control, and oxidative and nitrosative stress (Table 4).

The results obtained from this study revealed the genetic capacity of strain T4S, which suggests its bioprospecting in agriculture in enhancing plant growth. The predicted genes on the other hand may contribute to plant–bacterial interactions and ensure endosphere competence. Therefore, genome analysis of strain T4S provides information on the genes involved in sustainable plant growth and health.

**Sequence Data Information**
From the NCBI database output, the Bioproject is https://www.ncbi.nlm.nih.gov/bioproject/PRJNA706601; BioSample number is https://www.ncbi.nlm.nih.gov/biosample/SAMN 18138757 while the whole genome accession number is JAFNAY000000000.

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Author Contributions
All the mentioned authors contributed substantially and intellectually to this work. OOB designed the research, revised the work critically for important intellectual content, performed quality assurance, provided funding, project administration and resources. BSA was involved in data curation, formal analysis, investigation, visualization of data and writing of the original draft of the manuscript. ASA was involved in data curation, visualization of data, reviewing and thoroughly editing of the initial draft, validation and formal analysis.

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