

Draft Genome Sequence of Stenotrophomonas maltophilia Strain PE591, a Polyethylene-Degrading Bacterium Isolated from Savanna Soil

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ABSTRACT We report the genome sequence of a polyethylene-degrading bacterial strain identified as Stenotrophomonas maltophilia strain PE591, which was isolated from plastic debris found in savanna soil. The genome was assembled in 16 scaffolds with a length of 4,751,236 bp, a GC content of 66.5%, and 4,432 predicted genes.

Stenotrophomonas maltophilia strain PE591 was isolated from plastic debris found in
the soil of the Brazilian Cerrado biome ([1\)](#page-1-0). The strain showed both metabolic activity and cellular viability after incubation with unpretreated polyethylene (PE) (molecular weight, 191,000) films as the sole carbon source for periods of up to 90 days [\(1\)](#page-1-0). Moreover, S. maltophilia PE591 was capable of inducing significant physicochemical changes in PE after a 90-day incubation, revealing its great potential for plastic biodegradation processes [\(1\)](#page-1-0). The Stenotrophomonas genus currently comprises 20 species, and its first species, S. maltophilia, was described in 1993 [\(2](#page-1-1)[–](#page-1-2)[4](#page-1-3)). S. maltophilia is a Gram-negative, obligate aerobic, rodshaped, and motile bacterium that is considered to be an important human pathogen [\(5\)](#page-1-4). Stenotrophomonas species are ubiquitous microorganisms that colonize multiple natural (e.g., soils and plants) and clinical environments ([6,](#page-1-5) [7\)](#page-1-6). In addition, Stenotrophomonas spp. may have resistance to different metals and antibiotics; therefore, they qualify as promising microorganisms for bioremediation applications [\(6](#page-1-5)).

S. maltophilia strain PE591 was grown in 5ml of nutrient broth medium (Difco, Holland) and incubated for 24 h at 28°C with agitation (150 \times g). Subsequently, cells were harvested by centrifugation at 5,500 \times q for 5 min at 4°C. Bacterial DNA was purified using the GenElute bacterial genomic DNA kit (Sigma-Aldrich, USA). A sequencing library was created using the MiSeq reagent kit (Illumina, San Diego, CA) according to the manufacturer's instructions, the MiSeq system user guide, revision L (part number 15027617; Illumina), was used in the sequencing protocol, and genome sequencing was performed with the Illumina MiSeq system $(2 \times 300$ -bp paired-end reads) (Macrogen, Seoul, South Korea). The resultant reads were sub-jected to quality analysis using FastQC software v. 0.11.3 [\(8](#page-1-7)). Sequence reads were de novo assembled following the A5-miseq pipeline, which includes Trimmomatic v. 0.35 to trim lowquality sequences (Phred scores of $<$ 20) and IDBA-UD v. 1.1.1 to assemble contigs ([9](#page-1-8)[–](#page-1-9)[11\)](#page-1-10). The package Stats from BBmap v. 38.76 was used to generate assembly statistics ([12](#page-1-11)). The genome was annotated with Prokka v. 1.14.6 using the UniProt database [\(13\)](#page-1-12). The completeness and contamination of genomic data were estimated by CheckM v. 1.0.13 ([14](#page-1-13)). Coverage was assessed with SAMtools v. 1.9, Bowtie2 v. 2.3.4.1, and the package Pileup from BBmap v. 38.76 [\(12,](#page-1-11) [15](#page-2-0), [16](#page-2-1)). Default parameters were used for all software unless otherwise noted. Finally, the

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taxonomic classification was assessed following the analysis of (i) the average nucleotide identity (ANI) obtained through the web server JSpeciesWS [\(http://jspecies.ribohost.com/](http://jspecies.ribohost.com/jspeciesws/#analyse) [jspeciesws/#analyse](http://jspecies.ribohost.com/jspeciesws/#analyse)), which generates the ANIb (ANI algorithm using BLAST), and (ii) digital DNA-DNA hybridization (dDDH) obtained through the Type (Strain) Genome Server (TYGS) [\(https://tygs.dsmz.de\)](https://tygs.dsmz.de) ([17](#page-2-2), [18](#page-2-3)).

The genome sequencing generated a total of 13,411,628 reads, with a total of 4,019,163,037 bp. The genome of S. maltophilia PE591 was assembled in 16 contigs, with a total length of 4.75 Mb and a GC content of 66.5%. Genome coverage, completeness, and contamination were ~800 \times , 100%, and 0%, respectively. N₅₀ and L₅₀ values for the assembly were 473,413 bp and 3 contigs, respectively. Genomic functional features include a total of 4,432 genes, 4,350 protein-coding genes, 7 rRNAs, 74 tRNAs, and 1 transfer-messenger RNA.

The genomic ANI of S. maltophilia PE591 in relation to Stenotrophomonas maltophilia strain NCTC10258 (GenBank accession number [NZ_LS483377.1](https://www.ncbi.nlm.nih.gov/nuccore/NZ_LS483377.1)) was 97.75% (ANIb). The dDDH (d4) value provided by the TYGS against Stenotrophomonas maltophilia NBRC 14161 (RefSeq assembly accession number [GCF_001591205.1](https://www.ncbi.nlm.nih.gov/assembly/GCF_001591205.1/)) was 80.2%. According to these results and the minimum parameters for taxonomic classification of prokaryotes using genomic data ([19\)](#page-2-4), strain PE591 is classified as a novel strain of Stenotrophomonas maltophilia. Unraveling the genomes of PE degraders contributes to efforts to find strategies to address the worldwide accumulation of persistent plastics. The potential of S. maltophilia PE591 to break and to oxidize the most recalcitrant plastic qualifies it as a promising biological tool for tackling this urgent global issue.

Data availability. This whole-genome shotgun project has been deposited in DDBJ/ ENA/GenBank under the accession number [JAAOKO000000000.1,](https://www.ncbi.nlm.nih.gov/nuccore/JAAOKO000000000.1) with BioProject accession number [PRJNA609181](https://www.ncbi.nlm.nih.gov/bioproject/PRJNA609181) (sample, Stenotrophomonas sp. strain PE591 [BioSample accession number [SAMN14238446\]](https://www.ncbi.nlm.nih.gov/biosample/SAMN14238446); experiment, PE591 [SAR accession number [SRX10931469\]](https://www.ncbi.nlm.nih.gov/sra/SRX10931469); run, PE_591_1.fastq.gz [SRA accession number [SRR14584137\]](https://trace.ncbi.nlm.nih.gov/Traces/sra/?run=SRR14584137)).

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We declare no conflicts of interest regarding this article.

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