**Geminicoccus flavidas** sp. nov. and **Geminicoccus harenae** sp. nov., two IAA-producing novel rare bacterial species inhabiting desert biological soil crusts

Zhu-Ming Jiang1,2, Yang Deng1,2, Xue-Fei Han1,2, Jing Su1, Hao Wang1, Li-Yan Yu1 and Yu-Qin Zhang1,2*

1Institute of Medicinal Biotechnology, Chinese Academy of Medical Sciences, Beijing, China, 2State Key Laboratory of Dao-di Herb, Beijing, China

Two Gram-staining negative strains (CPCC 101082T and CPCC 101083T) were isolated from biological sandy soil crusts samples collected from Badain Jaran desert, China. Both isolates were heterotrophic phototroph, could produce indole-3-acetic acid. The 16S rRNA gene sequences of these two strains were closely related to the members of the family Geminicoccaceae, showing high similarities with Geminicoccus roseus DSM 18922T (96.9%) and Arboricoccus pini B29T1 (90.1%), respectively. In phylogenetic tree based on 16S rRNA gene sequences, strain CPCC 101082T and CPCC 101083T formed a robust distinct clade with Geminicoccus roseus DSM 18922T within the family Geminicoccaceae, which indicated that these two isolates could be classified into the genus Geminicoccus. The growth of strain CPCC 101082T occurred at 15–42°C and pH 4.0–10.0 (optima at 28–37°C and pH 6.0–8.0). The growth of strain CPCC 101083T occurred at 4–45°C and pH 4.0–10.0 (optima at 25–30°C and pH 6.0–8.0). The major cellular fatty acids of CPCC 101082T and CPCC 101083T contained C18:1ω7c/C18:1ω6c, cyclo-C19:0ω8c, and C16:0. Q-10 was detected as the sole respiratory quinone. Diphosphatidylglycerol, phosphatidylglycerol, phosphatidylcholine, phosphatidylethanolamine, an unidentified phospholipid and an unidentified aminolipid were tested in the polar lipids profile. The genomes of the two isolates were characterized as about 5.9 Mbp in size with the G+C content of nearly 68%. The IAA-producing encoding genes were predicated in both genomes. The values of average nucleotide identity were 80.6, 81.2 and 92.4% based on a pairwise comparison of the genomes of strains CPCC 101082T and CPCC 101083T and Geminicoccus roseus DSM 18922T, respectively. On the basis of the genotypic, chemotaxonomic and phenotypic characteristics, the strains CPCC 101082T (≡NBRC 113513T=KCTC 62853) and CPCC 101083T (≡NBRC 113514T=KCTC 62854) are proposed to represent two novel species of the genus Geminicoccus with the names Geminicoccus flavidas sp. nov. and Geminicoccus harenae sp. nov.

**KEYWORDS**

Geminicoccus flavidas, Geminicoccus harenae, average nucleotide identity, pan-genome, biological soil crusts
Highlights

This is the first report about the rare bacterial genus *Geminicoccus* was obtained from the desert niches. The primary goal of the present research was to identify the *Geminicoccus* cultures from Badain Jaran Desert, and to study the properties of *Geminicoccus* members inhabiting desert niches. The detailed phenotypic and genotypic properties resulted from these strains could be helpful to systematically demonstrate the ecological adaptation mechanism and their ecological function of the rare bacteria. As a result, strains CPCC 101082\(^1\) and CPCC 101083\(^2\), isolated from sandy soil samples collected from Badain Jaran Desert, showed indole-3-acetic acid (IAA)-producing activity. Based on the phenotypic and genotypic data, two novel species *Geminicoccus flavidas* sp. nov. and *Geminicoccus harenae* sp. nov., are proposed, with the isolates CPCC 101082\(^1\) and CPCC 101083\(^2\) as the type strains, respectively, for which the respective names *Geminicoccus flavidas* sp. nov and *Geminicoccus harenae* sp. nov. are proposed.

Introduction

The genus *Geminicoccus* was initially proposed by Foesel et al. (2007), with the type species *Geminicoccus roseus* (type strain DSM 18922\(^1\)) as the sole species. Originally, the genus *Geminicoccus* was classified in the order *Rhodospirillales* without being designed into any family. As of 2008, based on the core genes analysis on the available genomes of the members in the order *Rhodospirillales* and the newly isolated strain B29T\(^1\), which was isolated from the endophytic microbial community of a *Pinus pinaster* treetrunk, Proença et al. established the family *Geminicoccaceae*, with the genus *Geminicoccus* as the type genus, and also proposed a new genus *Arboricoccus*, with strain B29T\(^1\) being the type strain of the type species *Arboricoccus pini* (Proença et al., 2018). There are currently only four recognized genera: *Arboricoccus*, *Deflavicoccus*, *Geminicoccus*, and *Tistrella* being classified in the family *Geminicoccaceae* and there is one species in each genus. To summarize, currently, only strain DSM 18922\(^1\), isolated from a marine aquaculture biofilter, is accommodated in the genus *Geminicoccus*. Cells of the genus *Geminicoccus* are diplococci, non-motile, aerobic and heterotrophic phototroph. The present study aimed to establish the taxonomic status of two newly isolated *Geminicoccus* strains (CPCC 101082\(^1\) and CPCC 101083\(^2\)) recovered from desert biological soil crusts samples collected from Badain Jaran desert, China. The isolates were pairwise compared with one another and as well with the type strain DSM 18922\(^1\) of the currently described sole *Geminicoccus* species, and further studied using a polyphasic taxonomic approach. The isolates, strains CPCC 101082\(^1\) and CPCC 101083\(^2\), were found to represent two new *Geminicoccus* species, respectively, for which the respective names *Geminicoccus flavidas* sp. nov and *Geminicoccus harenae* sp. nov. are proposed.

Materials and methods

Acquisition of samples

The sandy soil samples were collected from Badain Jaran desert in Alxa Right Banner, Alxa League, China. Six samples sandy soil samples were collected at random from a hinterland area (39°34′24″N, 101°47′56″E, 1286 mH) characterized by cyanobacteria-dominated crusts, pooled and marked as BJDC. Another sandy soil mixture labeled as BJDM was obtained around a sand dune (39°18′45″N, 101°55′37″E, 1580 mH) with moss-dominated crusts. These samples were separately sealed in sterile envelopes and sent to the laboratory within 1 week after collection. All samples were immediately processed for isolation of microorganisms after arriving at the laboratory, and the remaining samples were maintained at −80°C.

Isolation and identification of bacterial strains

After dilution of the sample using 0.1% (w/v) NaCl solution mixed with 0.1% sodium pyrophosphate, 200 μL of 10⁻⁴ concentration soil-suspension was spread onto the isolation medium Difco™ Marine Agar 2,216, which contained (L⁻¹): 19.45 g NaCl, 8.8 g MgCl₂, 5 g peptone, 3.24 g Na₂SO₄, 1.8 g CaCl₂, 1 g yeast extract, 0.55 g KCl, 0.16 g NaHCO₃, 0.1 g ferric citrate, 0.08 g KBr, 34 mg SrCl₂, 22 mg Boric acid, 8 mg disodium phosphate, 4 mg sodium silicate, 2.4 mg NaF, 1.6 mg NH₄NO₃, 15 g agar; pH 7.6-7.8. Nystatin (50 mg L⁻¹) and potassium dichromate (30 mg L⁻¹; Shi et al., 2019) were added to the isolation medium to prevent the growth of fungi that might be present. After 4 weeks of incubation at 28°C, distinct colonies were picked and streaked onto PYG plates (5.0 g L⁻¹ yeast extract, 3.0 g L⁻¹ peptone, 10.0 mL L⁻¹ glycerol, 1.25 g L⁻¹ betaine hydrochloride, 1.25 g L⁻¹

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Abbreviations: DPG, Diphosphatidylglycerol; PG, Phosphatidylglycerol; PE, Phosphatidylethanolamine; PC, Phosphatidylcholine; PL, Unidentified phospholipid; AL, Unidentified aminophospholipid; ANI, Average nucleotide identity; dDDH, Digital DNA−DNA hybridization; IAA, Indole-3-acetic acid.
sodium pyruvate, 15 g L−1 agar) until the isolated and uniform colonies appeared. The purified culture was temporarily maintained on GYM agar (4 g glucose, 4 g yeast extract, 10 g malt extract, 2 g CaCO₃, 15 g agar, pH 7.0–7.2) at 4°C and stored as aqueous glycerol suspension (20%, v/v) at −80°C.

The reference strain of G. roseus DSM 18922¹ obtained from DSMZ was included in parts of the following assays as a parallel.

The newly isolated bacterial strains were primarily identified according to the 16S rRNA gene sequence comparison in the following steps.

Extraction of the genomic DNA of strains CPCC 101082² and CPCC 101083³ and sequencing of PCR amplification products of 16S rRNA genes were performed as described by Li et al. (2007). The obtained sequences were compared with available 16S rRNA gene sequences from GenBank using the BLAST program and EzBioCloud⁴ to determine the approximate phylogenetic affiliations of the strains (Kim et al., 2012). Multiple alignments with sequences of the most closely related taxa and calculations of levels of sequence similarities were carried out using MEGA version 11.0 (Tamura et al., 2021). Phylogenetic trees were inferred using the neighbor-joining method (Saitou and Nei, 1987) with K₂₀ values (Kimura, 1979; Kimura, 1980) and complete deletion gaps, and the maximum-parsimony (Kluge and Farris, 1969) and maximum-likelihood (Felsenstein, 1981) methods. The topology of the phylogenetic tree was evaluated by the bootstrap resampling method of Felsenstein (Felsenstein, 1985) with 1,000 replicates.

Observation of morphological properties and physiological tests

Cell Gram-staining was performed by standard Gram’s reaction. Cell morphology was observed by a light microscope and confirmed using transmission electron microscopy (JEM-1010, JEOL). The mobility of the cells was checked by inoculating colonies into GYM medium (g L−1; glucose 4, yeast extract 4, malt extract 10, calcium carbonate 2, pH 7.5) with 0.3% (w/v) agar and incubated at 30°C for 5 days.

The growth conditions of the isolates were tested using GYM, Tryptic soy agar (TSA, Difco), Reasoner’s 2A agar (R2A, Difco), nutrition agar (NA) and PYG agar by incubating the cultures for up to 7 days. The temperature range of cell growth was tested at 4, 10, 15, 20, 25, 28, 30, 32, 35, 37, 40, and 45°C using GYM slant medium in multi-thermo incubators (Eyela, MTI-20; Tokyo Rikakikai). The growth of the strains at different pH values (pH 4–12, at intervals of 1 pH unit) and NaCl concentrations (0–15%, w/v, at intervals of 1%) was examined using GYM broth at 30°C. Growth under anaerobic conditions was determined on GYM slant medium for 7 days at 30°C in an anaerobic incubator. Growth under light and dark conditions was monitored on GYM agar medium for 7 days at 30°C in an illumination incubator and ordinary constant temperature incubator, respectively. Cells were collected by centrifugation. About 0.5 g of the pellets were resuspended in 15 ml saline solution then centrifuged to remove the remained medium. Pigments were extracted by chloroform and analysed by HPLC (Reddy et al., 2006) and absorption spectra between 190 and 900 nm were recorded in a UV/Vis spectrophotometer (TU-1901, PERSEE; Kömpfer et al., 2003).

Catalase activity was evaluated by bubble production in 3% (v/v) H₂O₂, and oxidase activity was assessed by API oxidase reagent (bioMérieux). Metabolic characteristics were examined using Biolog GEN III (MicroPlate), API 50CH (bioMerieux), API 20NE (bioMérieux) and API ZYM test kits (bioMérieux) according to the manufacturer’s instructions. H₂S production and nitrate reduction were tested according to previously described procedures (Dong et al., 2014). Endoglucanase activity was examined using CMC- Na screening medium (Reinhold-Hurek et al., 1993). The abilities to hydrolyse gelatin and starch and other physiological tests were carried out as described by Gonzalez et al. (1978), Voges-Proskauer (VP) and Methyl-Red (MR) tests were performed on glucose-peptone broth medium. The ability of the strains to produce IAA was tested using colorimetric methods (Bric et al., 1991). Cells in the logarithmic phase of growth were inoculated into 1% tryptone aqueous solution containing 3 mmol/L L-tryptophan and then cultured at 30°C for 3 days. The concentration of IAA standard solution (0, 0.675, 1.25, 2.5, 5, 10, 20, 40, 50, 60, 80, 100 mg/L) was taken as the abscissa and the absorbance value of the corresponding concentration at 540 nm as the ordinate to draw a scatter diagram, and a linear trend line was added to obtain the IAA standard curve, as shown in Supplementary Figure S1.

Chemotaxonomic tests

Cells for chemotaxonomic and molecular systematic studies were collected from GYM broth at 30°C for 5–7 days, except the cells for fatty acids analysis. The polar lipids were extracted and isolated by two-dimensional TLC and identified by previously described procedures (Minnikin et al., 1984). Ubiquinone(s) was extracted and purified according to the method described previously and analyzed by HPLC (Collins et al., 1977). Cultures for cellular fatty acids extraction were harvested by using TSB broth (Difco) at 30°C for 5 days. Analysis of the whole-cell fatty acid profile was carried out by the Sherlock Microbial Identification System (MIDI) according to the manufacturer’s instructions (Kroppenstedt, 1985).

Whole-genome study

Genome sequencing, assembly and quality evaluation

Genome sequencing was performed on an Illumina HiSeq 4,000 system (illumina) at BGI company (Shenzhen, China). Genomic DNA was sheared randomly to construct three read libraries with lengths of 300 bp by a Bioruptor ultrasonicator (Diagenode, Denville, NJ, United States) and physico-chemical methods. The paired-end fragment libraries were sequenced.

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¹ http://www.ezbiocloud.net/
according to the Illumina HiSeq 4,000 system's protocol. Raw reads of low quality from paired-end sequencing (those with consecutive bases covered by fewer than five reads) were discarded. The sequenced reads were assembled using SOAPdenovo v1.05 software. The completeness and contamination of the genomes were accurately estimated by the CheckM pipeline.

Genomic component prediction and gene annotation

Gene prediction was performed on the genome assembly by glimmer35 with Hidden Markov models. The trRNAscan-SE (Lowe and Eddy, 1997), RNAmmer and the Rfam databases were employed for sorting trNA, rRNA and sRNAs, respectively. The annotation of tandem repeats was conducted using the Tandem Repeat Finder,4 and the minisatellite DNA and microsatellite DNA were selected based on the number and length of repeat units. The Genomic Island Suite of Tools (GIST) was used for genomic islands analysis8 with the IslandPath-DIOMB, SIGI-HMM and IslandPicker methods. Prophage regions were predicted using the PHAge Search Tool (PHAST) web server,9 and CRISPR identification was conducted using CRISPRFinder.

The best hit was abstracted using the Blast alignment tool for function annotation. Seven databases, KEGG (Kyoto Encyclopedia of Genes and Genomes), COG (Clusters of Orthologous Groups), NR (Non-Redundant Protein Database databases), Swiss-Prot, GO (Gene Ontology), TrEMBL and EggNOG, were used for general function annotation.

The protein sequences of each genome annotated by the abovementioned methods were used for downstream analysis such as stress-response gene retrieval and pan-genome analysis. For pathway analysis, the predicted protein sequences were uploaded to the KEGG Automatic Annotation Server using the “for prokaryotes” and “bidirectional best hit” options. The UniProt and Interpro10 websites were used for validation of the stress response genes. Predictions of gene clusters for natural products were performed using antiSMASH (Blin et al., 2019).

Whole genome-based taxonomy

The draft assembly of the genomes was submitted to GenBank to be publicly accessible. The draft assembly genomic sequence (accession number: ATYL00000000) of the strain Geminicoccus roseus DSM 189227 was deposited in the NCBI database. The genomic G + C content was calculated from the draft genome sequences. Genomic robust indexes, i.e., ANI (average nucleotide identity) and dDDH (digital DNA–DNA hybridisation) were calculated for the definition and classification of the species. The ANN Calculator9 (Yoon et al., 2017) and Genome-to-Genome Distance Calculator10 (Meier-Kolthoff et al., 2013) were used to calculate the ANI and dDDH, respectively.

The Bacterial Pan-genome Analysis (BPGA) pipeline was applied for analysis of the genomic diversity of the Geminicoccus members. The protein sequences of the strains CPCC 1010828 and CPCC 1010839 were predicted by glimmer 3.02, and the protein sequence of the strain G. roseus DSM 189227 was annotated by RAST 2.0. Pan-genome analysis was performed by BPGA 1.3 using default settings (Chaudhari et al., 2016). A total of three protein sequence files annotated from the three corresponding strains’ whole genome sequences were used to generate orthologous gene/protein clusters (homologous families) using the USEARCH clustering tool and to construct the phylogenetic tree using the concatenated core genes using BPGA.

Each homologous family was given a conserved value based on the frequency in the three genomes. Here, we defined the homologous gene families conservation value (CV) as 3, when a certain homologous gene was common shared by all of the type strains of these three species. Different CVs reflect the distribution frequency of the homologous gene among the three strains. A higher CV indicates that the gene is more widely distributed in the strains of the genus Geminicoccus and the gene family is more conserved in the genus Geminicoccus. In the pan-genome of the three strains, if the CVs of some homologous families were 3, these homologous families were considered part of the core genome; homologous families with CVs of 2 or 1 were regarded as accessory genes or unique genes, respectively. The core, accessory, unique and exclusively absent genes were retrieved using the USEARCH clustering tool. BPGA was used to perform the evolutionary analysis based on concatenated core gene alignments and the binary (presence/absence) pan-matrix. The gene matrix was calculated using similarity or dissimilarity in the contribution of genes to orthologous gene clusters. For the core genome-based phylogenetic tree, BPGA first extracted the protein sequences (excluding paralogs) from 20 random orthologous gene clusters to generate the core genome phylogeny tree. BPGA automated multiple sequence alignments using MUSCLE. All alignments were concatenated, and a neighbor-joining phylogenetic tree was constructed.

Results and analysis

Isolation and primary taxonomic study results on Geminicoccus strains

Strains CPCC 1010827 and CPCC 1010839 were recovered from sandy soil samples BJDC and BJDM, respectively. The almost full-length 16S rRNA gene sequences (1,449 and 1,453 bp) of strains CPCC 1010827 and CPCC 1010839 were

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5 http://www.ccbcb.umd.edu/software/glimmer/
4 http://tandem.bu.edu/trf/trf.html
5 http://www5.esu.edu/cpsc/bioinfo/software/GIST/
6 http://phast.wishartlab.com/
7 https://www.uniprot.org/
8 https://www.ebi.ac.uk/interpro/
obtained. The comparison of the 16S rRNA gene sequences indicated that strains CPCC 101082<sup>T</sup> and CPCC 101083<sup>T</sup> were members of the family Geminicoccaceae, with the highest similarities to G. roseus DSM 18922<sup>T</sup> (CPCC 101082<sup>T</sup>: 96.9%; CPCC 101083<sup>T</sup>: 96.9%) and Arboricoccus pini B29T<sup>T</sup> (CPCC 101082<sup>T</sup>: 90.8%; CPCC 101083<sup>T</sup>: 90.1%). The similarity of the 16S rRNA gene sequences between CPCC 101082<sup>T</sup> and CPCC 101083<sup>T</sup> was 98.6%. These values were all lower than the similarity of 98.65%, which was proposed by Kim et al. as the threshold for differentiating two species (Kim et al., 2014). In the phylogenetic tree based on the 16S rRNA gene sequences of all genera within the order Rhodospirillales, strains CPCC 101082<sup>T</sup> and CPCC 101083<sup>T</sup> fell in the genus Geminicoccus lineage and formed a robust unique cluster with Geminicoccus roseus DSM 18922<sup>T</sup> in the neighbor-joining tree (Figure 1), which showed almost the same case in the maximum-parsimony tree and maximum-likelihood tree. These results suggested that strains CPCC 101082<sup>T</sup> and CPCC 101083<sup>T</sup> represent two novel species of the genus Geminicoccus.

### Phenotypic properties

**Morphological and physiological characteristics**

Strains CPCC 101082<sup>T</sup> and CPCC 101083<sup>T</sup> grew well on GYM agar. Better growth was observed under light compared with dark conditions. White/light yellow colonies with a wrinkled surface and a maximum diameter of 3–5 mm with irregular edges on GYM were observed after incubation in an illumination incubator for 8 days at 30°C. Strain CPCC 101083<sup>T</sup> formed pale pink colonies with a slightly wrinkled surface and a maximum diameter of 1–3 mm with regular edges on GYM after incubation in an illumination incubator for 8 days at 30°C. The cells of strains CPCC 101082<sup>T</sup> and CPCC 101083<sup>T</sup> were gram-negative, coccoid or short-rods. Both strains were aerobic. The absorption spectra of pigments extracted from the cells of strain CPCC 101083<sup>T</sup> exhibited the same as of carotenoids, with major absorption maxima at 467, 498 and 525 nm.

The growth of the strain CPCC 101082<sup>T</sup> occurred at 15–42°C (optimum 28–37°C), and pH 4.0–10.0 (optimum pH 6.0–8.0). The range of NaCl tolerance was 0–4% (w/v). Strain CPCC 101083<sup>T</sup> grew at 4–45°C (optimum 25–30°C) and pH 4.0–10.0 (optimum pH 6.0–8.0). The range of NaCl for growth was 0–5% (w/v).

In the fermentation broth of strains CPCC 101082<sup>T</sup> and CPCC 101083<sup>T</sup>, 3-indole-acetic acid (IAA) was detected. As shown in Supplementary Figure S1, the linear regression equation \( y = 0.0244x + 0.0482, r^2 = 0.9998 \), had a good fit. Accordingly, the IAA content produced by the test strains could be calculated according to this equation. The IAA content yielded from the fermentation broth of CPCC 101082<sup>T</sup>, CPCC 101083<sup>T</sup> and DSM 18922<sup>T</sup> was 3.83 ± 0.02, 4.75 ± 0.09, and 2.10 ± 0.05 mg/L, respectively.

**FIGURE 1**

Neighbour-joining tree based on 16S rRNA gene sequences showing the relationship of the strains CPCC 101082<sup>T</sup>, CPCC 101083<sup>T</sup> and Geminicoccus roseus DSM 18922<sup>T</sup> with representatives of the order Rhodospirillales. Filled circles indicate that the corresponding nodes were also recovered in the trees generated with the maximum-likelihood and maximum-parsimony methods. Bootstrap values (>50 %) are shown as percentages of 1000 replicates. Bar, 2 nt substitutions per 100 nt.
The activity of alkaline phosphatase of strain CPCC 101082\(^2\) and CPCC 101082\(^1\) was positive. The endoglucanase activity of strain CPCC 101082\(^1\) was positive, while the endoglucanase activity of strain CPCC 101083\(^1\) was weakly positive. Strain CPCC 101082\(^2\) could hydrolyze urea, while CPCC 101083\(^2\) could not. The activities of nitrate reduction, oxidase and catalase of strain CPCC 101083\(^2\) were positive, but those of strain CPCC 101082\(^2\) were negative. Other detailed physiological characteristics and the differential phenotypic characteristics of strains CPCC 101082\(^2\) and CPCC 101083\(^2\) with respect to the closest phylogenetic neighbor G. roseus DSM 18922\(^1\) and other phylogenetic neighbors are shown in Table 1.

### Chemotaxonomic characteristics

The polar lipids of strains CPCC 101082\(^1\) and CPCC 101083\(^1\) mainly consisted of diphasophatidylglycerol (DPG) phosphatidylglycerol (PG), phosphatidylcholine (PC), phosphatidylethanolamine (PE), an unidentified phospholipid (PL) and an unidentified aminolipid (AL) (Supplementary Figure S2). The sole respiratory quinone was detected as Q-10. The fatty acids (>5%) of strain CPCC 101082\(^2\) contained cyclo-C\(_{18:0}\)Δ9c (40.4%), Summed Feature 8 (C\(_{16:0}\)ω7c/ C\(_{16:0}\)ω6c; 29.0%) and C\(_{16:1}\)ω6c (10.8%) with minor (1–5%) amounts of C\(_{17:0}\)ω6c (4.5%), Sum In Feature 4 (iso-C\(_{17:0}\)) I and/or anteiso-C\(_{17:1}\) B; 2.2%), Sum In Feature 3 (C\(_{16:0}\)ω7c and/or iso-C\(_{15:0}\)2-OH; 2.1%), C\(_{18:0}\)ω5c (2.0%), C\(_{18:1}\)ω9c (1.8%) and C\(_{18:3}\)-OH (1.8%). The major fatty acids (>5%) of strain CPCC 101083\(^2\) were Summed Feature 8 (C\(_{16:0}\)ω7c/C\(_{16:0}\)ω6c; 37.4%), cyclo-C\(_{16:0}\)Δ9c (31.8%), C\(_{16:1}\)ω6c (14.1%) and C\(_{18:3}\) (8.8%), with minor (1–5%) amounts of C\(_{18:3}\)-3-OH (1.8%); Supplementary Table S1).

The profile of the chemotaxonomic properties stated above well supported the classification of strains CPCC 101082\(^1\) and CPCC 101083\(^1\) as two members of the genus Geminicoccus. However, strains CPCC 101082\(^1\) and CPCC 101083\(^1\) differed from each other and as well with the only validly identified Geminicoccus member G. roseus DSM 18922\(^1\) in some physiological properties and the cellular fatty acid profile (Table 1; Supplementary Table S1).

Filled circles indicate that the corresponding nodes were also recovered in the trees generated with the maximum-likelihood and the maximum-parsimony methods. Bootstrap values (>50%) are shown as percentages of 1,000 replicates. Bar, 2 nt substitutions per 100 nt.

### Genome properties

Genome sequencing of strain CPCC 101082\(^2\) yielded a draft genome containing 5,871,762 bp, assembled from 1,148 qualified scaffolds, with a coverage of 384-fold and an N50 length of 180,117 bp. The G + C content in the genomic DNA of strains CPCC 101082\(^2\) and CPCC 101083\(^2\) was calculated as 68.2 and

| Characteristic | CPCC 101082\(^2\) | CPCC 101083\(^2\) | DSM 18922\(^1\) |
|----------------|------------------|------------------|------------------|
| Colony color   | white to light yellow | pale pink | whitish-grey to light pink |
| Cell morphology| coccoid to rod | coccoid to short rod | coccoid |
| Temperature range for growth (°C; Optimum) | 15–42 (28–37) | 4–45 (25–35) | 15–45 (30–35) |
| pH range for growth | 4.0–10.0 (6.0–8.0) | 4.0–10.0 (6.0–8.0) | 5.5–11.0 (6.0–8.0) |
| Tolerance of NaCl (%) | 0–4 | 0–5 | 0–10 |

### Hydrolysis of

- Leucine arylamidase
- Valine arylamidase
- Cystine arylamidase
- Trypsin
- α-chymotrypsin
- Acid phosphatase
- β-glucuronidase

### Utilization of (Biolog GEN III):

- Dextrin
- D-Maltose
- D-Cellbiose
- Gentiobiose
- α-D-Lactose
- D-Melibiose
- N-Acetyl-D-Glucosamine
- α-D-Glucose
- D-Mannose
- D-Galactose
- α-Methyl-Glucose
- L-Rhamnose
- D-Sorbitol
- D-Mannitol
- D-Arabinol
- N-Acetylglucosamine
- α-d-Glucose
- D-Glucosamine
- D-Glucuronic Acid
- L-Lactic Acid
- α-Keto-Glutaric Acid
- D-Malic Acid
- L-Malic Acid

(Continued)
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At least 63% of those are Gram-reaction-negative, positive for hydrolysis of starch and production of indole-3-acetic acid, negative for production of indole and His VP and MR tests. In API ZYM kits, all strains are positive for alkaline phosphatase, esterase (C₅), leucine arylamidase, and naphthol-AS-B1-phosphohydrolase, negative for lipase (C₁₆). α-galactosidase, β-galactosidase, α-glucosidase, β-glucosidase, N-acetyl-β-D-glucosaminidase, N-mannosidase and β-fucosidase. In Biolog GEN III (MicroPlate), all strains are positive for D-fructose, D-fucose, D-glucose-6-P, D-fructose-6-P, D-galacturonic acid, L-galactonic acid lactone, gluconolactone and acetic acid, are negative for D-trehalose, sucrose, D-turanose, stachyose, D-raffinose, β-methyl-D-glucoside, D-salicin, N-acetyl-β-D-mannosamine, N-acetyl-D-galactosamine, N-acetyl-neuraminic acid, inosine, glycerol, D-aspartic acid, D-serine, gelatin, glycol-L-proline, L-alanine, L-arginine, L-aspartic acid, L-glutamic acid, L-proline, glutamine, aspartic acid, p-hydroxy-phenylacetic acid, methyl pyruvate, D-lactic acid methyl ester, citric acid, bromo-succinic acid, α-aminobutyric acid, α-hydroxy-butyric acid, α-keto-butyric acid and propionic acid. All data were from this study, +, positive; −, negative; w, weakly positive.

67.9%, respectively, from the genome sequence. The DDDH values among strains CPCC 101082, CPCC 101083, and Geminicoccus roseus DSM 18922 ranged from 23.0 to 47.0%, far below the cut-off value (70%) used to classify bacterial strains of the same species (Audh et al., 2010); the ANI values among CPCC 101082, CPCC 101083 and Geminicoccus roseus DSM 18922 ranged from 90.6 to 92.4%, also much lower than the threshold for bacterial species delineation (95–96%; Kim et al., 2014; Supplementary Table S2).

The whole genome of strains CPCC 101082 and CPCC 101083 contained 5,733 and 5,596 genes, respectively (Table 2). The strain G. roseus DSM 18922, which was reported to be isolated from the biofilter of a recirculating marine aquaculture system composed of 5,260 genes, was included as a reference. The detailed genomic characteristics of these strains are summarized in Table 2. In the genomes of strains CPCC 101082, CPCC 101083 and G. roseus DSM 18922, the putatively encoding genes for superoxide dismutase (sodA and sodB), catalase (katA and katE), peroxidase (isoA) and UV radiation-resistant genes (uvrA, uvrB, uvrC, and uvrD) were retrieved, which might endow these strains the ability to adapt to the strong solar radiation in the desert. The DNA repair-related genes (recA, recD, recF, recG, recJ, recN, recO, recQ, recR, and recX), heat shock-related genes (dnaJ, dnaK, groEL, groES, groL, grpE, hrcA, hslR, hslU, hspB, hscC, hspD, hspI, hspQ, htpG, and htpX), cold shock-related genes (cspA), carbon assimilation-related genes (coxS, cutL, and cutM), carbon source starvation response-related genes (crsA) and osmotic stress-related genes (otsA, otsB, treS, treY, treZ, opuAA, opuAC, and betA) were also retrieved. These genes mentioned above were related to the microorganisms response to the abiotic-stress in extreme desert environment. The metabolism related genes were retrieved, such as carotenoid biosynthesis-related genes (crtA, b, c), in addition, genes encoding flagella (flgB, flgC, flgD, flgE, flgF, flgG, flgH, flgI, flgK, flaA, flaB, flaF, flaI, flaJ, flgC, flaL, flaM, flaN, flaP, flaQ, flaR, flaS, motA, motB, and cheL), arsenic resistance-related genes (arsB, arsC, and arsH), heavy metal ion resistance-related genes (copA, copB, copZ, cadA, thi, ccrR, ccdD, dnanG, trxA, and trxB), and multidrug resistance and efflux pump-related genes (acrA, acrB, b, bpeA, mdtA, mdtB, and mgeX) were also retrieved from the genomes of strains CPCC 101082, CPCC 101083 and G. roseus DSM 18922 (Figure 2).
Panel analysis of the genus *Geminicoccus*

A total of 17,468 protein-coding genes (Table 3) were sorted from the genomes of these three strains of the genus *Geminicoccus*, which were divided into 7,856 homologous families by cluster analysis. Histograms were constructed according to different CVs (Figure 3A). Among them, there were a total of 3,472 core genes commonly shared by these three strains (CV = 3), accounting for about 44.2% of the total number of homologous gene families. The accessory genes (1,334 genes; CV = 2) accounted for about 17.0% of the homologous gene families in the newly proposed species. The proportion of the unique genes (3,050 genes; CV = 1) was about 38.8%.

The relationship between the pan-genome size and the number of genomes of the genus and the relationship between the number of core genes and the number of genomes were deduced (Figure 3B) using all of the protein sequences extracted from these three strains of the genus *Geminicoccus*. The functional relationship between the pan-genome size ($f_{\text{pan}}$) and the number of genomes ($n$) was obtained by fitting, as follows:

$$f_{\text{pan}}(n) = 5379.93 \times n^{0.345402}$$

The functional relationship between the number of core genes ($f_{\text{core}}$) and the number of genomes ($n$) was obtained by fitting, as follows:

$$f_{\text{core}}(n) = 6908.74 \times c^{-0.262635n}$$

It could be inferred from the pan-genome fitting curve in Figure 3B that, with the increasing number of sequenced genomes, the pan-genome size became larger and larger, instead of tending to a plateau. Accordingly, it could be proposed that the pan-genome of the genus *Geminicoccus* is the open type, which suggests that the genus *Geminicoccus* has a strong ability to accept horizontal gene transfers.

Out of 7,856 genes/gene clusters, BPGA mapped 3,911 (49.8%) to KEGG (Kyoto Encyclopedia of Genes and Genomes) pathways, i.e., core gene (2,860, 73.1%), accessory genes (437, 11.2%) and unique genes (614, 15.7%). After filtering KEGG pathways related to eukaryotes, we obtained an overview of the metabolic pathway corresponding to the gene(s) in the pan-genome of the genus *Geminicoccus*. A large number of core genes (2,635) were involved in carbohydrate metabolism (15.8%), amino acid metabolism (13.2%), membrane transport (12.4%),...
TABLE 3 The pan-genome profile information of the genus Geminicoccus.

| Genome number | Organism name | No. of core genes | No. of accessory genes | No. of unique genes | No. of exclusively absent genes |
|---------------|---------------|------------------|-----------------------|---------------------|---------------------------------|
| 1             | DSM 18922    | 3,472            | 316                   | 1,421               | 1,018                           |
| 2             | CPCC 101082  | 3,472            | 1,187                 | 941                 | 147                             |
| 3             | CPCC 101083  | 3,472            | 1,165                 | 688                 | 169                             |

some other elementary metabolism (biosynthesis of amino acids, 5.1%; carbon metabolism, 4.0%; fatty acid metabolism, 1.4%; 2-oxoacrylic acid metabolism, 1.4% and degradation of aromatic compounds, 0.4%; 12.3%), energy metabolism (6.3%), metabolism of cofactors and vitamins (6%), nucleotide metabolism (4.6%), lipid metabolism (4.0%), xenobiotic biodegradation and metabolism (3.4%), translation (3.2%), metabolism of other amino acids (2.9%), signal transduction (2.7%), replication and repair (2.6%), biosynthesis of other secondary metabolites (1.9%), folding, sorting and degradation (1.9%), cell motility (1.8%), metabolism of terpenoids and polyketides (1.6%), drug resistance (1.4%), glycan biosynthesis and metabolism (1.3%). Accessory and unique genes appeared to be enriched in membrane transport, carbohydrate metabolism, amino acid metabolism, as well as energy metabolism. Among the accessory genes (368), the major portion of genes were related to membrane transport (19.0%), carbohydrate metabolism (13.6%), some other elementary metabolism (biosynthesis of amino acids, 3.0%; carbon metabolism, 2.8%; fatty acid metabolism, 2.4%; degradation of aromatic compounds, 0.8%; 10.1%), amino acid metabolism (10.1%), signal transduction (6.5%), energy metabolism (6.5%), lipid metabolism (5.4%), metabolism of cofactors and vitamins (5.4%), xenobiotics biodegradation and metabolism (4.4%), nucleotide metabolism (3.8%), metabolism of other amino acids (3.5%), cell motility (3.3%), metabolism of terpenoids and polyketides (2.5%), replication and repair (1.9%), biosynthesis of other secondary metabolites (1.6%) and drug resistance (1.1%). Unique genes (576) seemed to be mainly enriched in carbohydrate metabolism (15.5%), membrane transport (15.1%), amino acid metabolism (12.2%), energy metabolism (11.3%), xenobiotics biodegradation and metabolism (6.4%), lipid metabolism (5.7%), signal transduction (5.6%), metabolism of cofactors and vitamins (4.9%), nucleotide metabolism (4.0%), biosynthesis of other secondary metabolites (2.6%), metabolism of terpenoids and polyketides (2.3%) and the metabolism of other amino acids (1.7%; Figure 4).

In the phylogenetic trees based on the concatenated core genes and binary gene presence/absence matrix (pan-matrix), strains CPCC 101082 and CPCC 101083 clustered in a unique branch in the lineage of the genus Geminicoccus, with the closest evolutionary distance with the branch formed by G. roseus DSM 18922T (Supplementary Figures S3, S4).

**Speculation of metabolic characteristics**

Two KEGG pathways for autotrophic carbon dioxide fixation of strains CPCC 101082 and CPCC 101083 were predicted (Supplementary Figure S5). The KEGG pathway diagram showed that the genes of strains CPCC 101082 and CPCC 101083 encoded some enzymes in autotrophic carbon dioxide fixation pathways in prokaryotes other than in the reductive pentose phosphate pathway (Supplementary Figures S5A,C). The KEGG pathway diagram also showed that the genes of strains CPCC 101082 and CPCC 101083 encoded some enzymes in the reductive pentose phosphate pathway of autotrophic carbon dioxide fixation (Supplementary Figures S5B,D). In the predicted KEGG pathway of the strain CPCC 101082 and CPCC 101083, some multidrug resistance and efflux pump related coding genes were retrieved, for example, acrA and acrB (multidrug efflux pump subunit AcrA and AcrB coding gene; Wang et al., 2017), bepF (Martin et al., 2009; efflux pump periplasmic linker BepF coding gene, may contribute to resistance to some drugs, such as sodium dodecyl sulfate and nalidixic acid), bpeA (multidrug efflux periplasmic linker protein BpeA coding gene), mdtA (Nagakubo et al., 2002) and mdtE (multidrug resistance protein MdtA and MdtE coding gene) and mexG (Aires et al., 1999) multidrug efflux RND transporter inhibitory subunit MexG coding gene).

**Taxonomic study conclusion**

The phenotypic and genotypic data supported the accommodation of strains CPCC 101082 and CPCC 101083 in the genus Geminicoccus and also distinguished these two strains from each other as well from other species of the genus Geminicoccus. Accordingly, we propose that strains CPCC 101082 and CPCC 101083 be classified as representatives of two new species of the genus Geminicoccus, with the names Geminicoccus flavidas sp. nov. and Geminicoccus harenae sp. nov.

**Description of Geminicoccus flavidas sp. nov.**

Geminicoccus flavidas (fla’vi.da. L. fem. adj. flavida yellowish).

Cells are Gram-staining-negative, cocoid to rods, non-motile and aerobic. Grows well on GYM agar and nutrient agar. Colonies on GYM agar are white/light yellow colonies with wrinkled, circular, convex, and opaque, approximately 1 mm in diameter after 5 days at 30°C (pH 7.0). Grows at 15–42°C and pH 4.0–10.0, with the optimum at 28–32°C and pH 6.0–8.0, respectively. Grows in NaCl at concentrations up to not more than 4%. Can utilize 3-methyl glucose, acetic acid, D-cellulbiose, dextrin, D-fructose, D-fructose-6-PO₄, D-fucose, D-galactose, D-galacturonic acid, D-glucose-6-PO₄, D-glucuronic acid, D-maltose, D-mannose, D-melibiose, formic acid, gentiobiose, glucuronamide, L-fucose, L-galactonic acid lactone, L-rhamnose, tween 40, α-D-glucose and
α-D-lactose as the sole carbon source, and D-arabinose, D-arabitol, D-ardonitol, D-fructose, D-fucose, D-galactose, D-glucose, D-lyxose, D-mannitol, D-mannose, D-ribose, D-sorbitol, D-tagatose, dulcitol, D-xylose, esculin ferric citrate, inositol, L-arabinose, L-arabitol, L-fucose, L-rhamnose, L-sorbose, L-xylose, potassium 5-keto-gluconate and xylitol can
be assimilated to produce acid. Positive for alkaline phosphatase, cystine arylamidase, esterase (C4), esterase lipase (C8), leucine arylamidase, naphthol-AS-BI-phosphohydrolase, trypsin and valine arylamidase, weakly positive for α-chymotrypsin in API ZYM strip. The cellular polar lipid system includes diphasosphatidylglycerol (DPG), phosphatidylglycerol (PG), phosphatidylcholine (PC), phosphatidylethanolamine (PE), an unidentified phospholipid (PL) and an unidentified aminolipid (AL). The sole respiratory quinone is Q-10. The major cellular fatty acids are C16:0, C18:0, C16:1ω7c, C18:1ω6c, and C18:1ω9c. The genome of the type strain is characterized by a size of 5.95 Mbp and the G + C content of 68.2%. The type strain CPCC 101082 (=NBRC 113513 = KCTC 62853°) was isolated from a hinterland area characterized by cyanobacteria-dominated crusts collected from Badain Jaran desert, China. The DDBJ/EMBL/GenBank accession numbers of 16S rRNA gene sequence and draft genome sequence of the strain CPCC 101083T are MK392027 and JABGCL000000000, respectively.

**Description of Geminicoccus harenae sp. nov.**

*Geminicoccus harenae* (ha.re’nae. L. gen. n. harenae of sand, of a desert, referring to the isolation source of the type strain from desert sand).

Cells are Gram-reaction-negative, coccoid to short-rods, non-motile and aerobic. Grows well on GYM agar and nutrient agar. Colonies on GYM agar are wrinkled, circular, convex, and opaque and opaque with a pale pink color, approximately 1 mm in diameter after 5 days at 30°C (pH 7.0). Grows at 4–45°C and pH 4.0–10.0, with the optimum at 25–30°C and pH 6.0–8.0. NaCl is not necessary for growth, while NaCl tolerance is 5.0% (w/v). Can utilize 3-methyl glucose, acetic acid, acetic acid, D-arabitol, D-cellobiose, dextrin, D-fructose, D-fructose-6-PO4, D-fucose, D-galactose, D-galacturonic acid, D-glucuronic acid, D-glucose-6-PO4, D-mannitol, D-mannose, D-sorbitol, formic acid, glucuronamide, L-fucose, L-galactonic acid lactone, L-histidine, L-lactic acid, L-rhamnose, myo-inositol, α-D-glucose and β-hydroxy-D,L-butyric acid as the sole carbon source, and amygdalin, arbutin, D-arabinose, D-arabitol, D-arabinitol, D-cellobiose, D-fructose, D-fucose, D-galactose, D-glucose, D-lyxose, D-mannose, D-rhamnose, D-sorbitol, D-tagatose, D-trehalose, dulcitol, D-xylose, esculin ferric citrate, inositol, L-arabinose, L-fucose, L-rhamnose, L-sorbose, L-xylose, N-acetyl-glucosamine, potassium 2-ketogluconate, potassium 5-ketogluconate, salicin and xylitol can be assimilated to produce acid. Positive alkaline phosphatase, cystine arylamidase, esterase(C4), esterase lipase(C8), leucine arylamidase, naphthol-AS-BI-phosphohydrolase, valine arylamidase and β-glucuronidase, weakly positive for acid phosphatase and trypsin in API ZYM strip. The predominant polar lipids are diphasosphatidylglycerol (DPG), phosphatidylglycerol (PG), phosphatidylcholine (PC), phosphatidylethanolamine (PE), an unidentified phospholipid (PL) and an unidentified aminolipid (AL). The sole respiratory quinone is Q-10. The major cellular fatty acids are C16:0, C18:0, C16:1ω7c, C18:1ω6c, and C18:1ω9c. The genome sequence is characterized by a size of 6.02 Mbp and the G + C content of 67.3%. The type strain CPCC 101083T (=NBRC 113514° = KCTC 62853°) was isolated from a sandy dune sample with moss-dominated crusts collected from Badain Jaran desert, China. The DDBJ/EMBL/GenBank accession numbers of 16S rRNA gene sequence and draft genome sequence of the strain CPCC 101083T are MK392027 and JABGCL000000000, respectively.

**Discussion**

Based on the genomic information, we summarized the genetic characteristics of the novel species *G. flavidas* sp. nov. and *G. harenae* sp. nov. accommodating the type strain recovered from a desert. The detailed phenotypic properties illustrated the abilities of these strains to adapt to harsh environmental stress. The genome-scale analysis of strains CPCC 101082 and CPCC 101083 indicated that adaptation of these organisms to desert niches was achieved through various response strategies to UV radiation, carbon starvation, desiccation, osmotic stress and so on.

Dehydration of bacterial cells due to desiccation caused severe damage to enzymes and electron transport chains, leading to the accumulation of free radicals (Billi and Potts, 2002), which in turn seriously damaged DNA (Dose et al., 1992). It is proposed that extreme radiation resistance in desert microorganisms is functionally related to the photochemical oxidation of desiccated cells. In such environments, aerobic organisms have antioxidant enzymes, such as superoxide dismutase, catalase and peroxidase, and some UV-radiation resistant related genes that are helpful in protecting metabolically active cells against endogenous reactive oxygen species generated from aerobic respiration. In the core genome of the genus *Geminicoccus*, 68 gene clusters (2.6%) were mapped to replication and repair (KO09124, a KEGG sub-category), consisting of DNA replication (ko03030), base excision repair (ko03410), nucleotide excision repair (ko03420), mismatch repair (ko03430), and homologous recombination (ko03440) et al. In the annotation results, some oxidative stress related genes, such as superoxide dismutase coding genes (sodA and sodB), thioredoxin/glutathione peroxidase (btuE), catalase coding genes (katA and katE), some UV-radiation resistant genes, such as the UvrABC repair system coding genes (uvrA, uvrB, and uvrC), DNA helicase II coding gene (uvrD) and some DNA repair related genes (recA, recD, recF, recG, recJ, recN, recO, recQ, recR, and recX) were retrieved (Figure 2).

In addition to mediating plant-microbe interaction, recent studies have shown that IAA conferred protection against environmental stresses such as UV radiation, salt, acidity, osmotic shock and heat shock (Bianco et al., 2009; Duca et al., 2014). The increasing trehalose, LPS (lipopolysaccharide), EPS (exopolysaccharide) and biofilm content induced by IAA
enhanced the resistance of bacteria to environmental stress (Bianco et al., 2009). Therefore, these novel and rare IAA-producing bacterial species in desert habitats, may promote the colonization of other dominant bacteria in desert habitats. The phenotypic assays confirmed that all the strains CPCC 101082T, CPCC 101083T and DSM 18922T could produce IAA. In the genomic category, the aldehyde dehydrogenase (EC 1.2.1.3) coding genes and amine oxidase coding gene (maoI) were retrieved from the genomes of CPCC 101082T, CPCC 101083T and G. roseu DSM 18922T. In addition, according to the distribution of IAA-producing genes among the three strains, the pathway of IAA production (Figure 5) was predicted through the annotation of the KEGG database.

The strain CPCC 101082T could hydrolyze urea, but CPCC 101083T could not. The difference in the copy numbers of urease-encoding genes might be the key factor leading to the difference in urea hydrolysis activity between the two strains. The carotenoid biosynthesis-related gene (crtrB) was retrieved from the genome of strain CPCC 101083T, not CPCC 101082T, which could help us understand that the methanol/acetone extracts from CPCC 101083T displaying absorbance maxima (418, 480 and 537 nm) were indicative of carotenoids other than those from strain CPCC 101082T. Bacterial secondary metabolism is a rich source of novel bioactive compounds with potential medicinal value. To find new drug-producing candidates, microbiologists are increasingly employing genome sequencing of a wide variety of microbes. Here,

\[ \text{FIGURE 5} \]
Putative overview of indole-3-acetic acid-producing pathway in the tryptophan metabolism pathways of the strain CPCC 101082T, CPCC 101083T, and Geminicoccus roseu DSM 18922T. Squares colored with green, blue and red represent the strain CPCC 101082T, CPCC 101083T, and DSM 18922T, respectively.
we identified biosynthetic sites by antiSMASH (antibiotic & Secondary Metabolite Analysis Shell). However, the results from the antiSMASH database showed that in the two strains, G. flavidas sp. nov. and G. harenae sp. nov., only 10 secondary metabolite gene clusters with moderate similarities to previously described secondary metabolite biosynthetic gene clusters were retrieved (Supplementary Table S3). These gene clusters exhibited 5–32% similarities to previously reported secondary metabolite biosynthetic gene clusters, such as bacillibactin NRP, ibomycin (Polyketide), NRP: cyclic depsipeptide + polyketide: Modular type I, oxalomycin B, alkaloid gene clusters and other unidentified secondary metabolite clusters attributable to NAPAA, terpene, thioamidites, RRE-containing and T1PKS types. Abundant functional genes (for instance, stress-responding genes) were found in the core genomes of the newly proposed species, G. flavidas and G. harenae, which are assumed to contribute greatly to their survival abilities in harsh desert environments. Therefore, from an evolutionary perspective, the ability to endure harsh environments might endow the bacteria with advantages in coping with desert environments; being the indispensable constituents of the pioneer biological population in the desert environment, these microorganisms contributed greatly to the development of the biosphere.

Data availability statement

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found in the article/Supplementary material.

Author contributions

Z-MJ, YD, and X-FH carried out the experiments and prepared the manuscript, JS, HW, and L-YY collected the samples. Z-MJ and Y-QZ designed the research and analyzed the data. All authors contributed to the article and approved the submitted version.

Funding

This research was supported by CAMS Innovation Fund for Medical Sciences (CIFMS, 2021-12M-1-055), National Natural Science Foundation of China (32170021), Beijing Natural Science Foundation (5212018), Key project at central government level-the ability establishment of sustainable use for valuable Chinese medicine resources (2060302), and the National Infrastructure of Microbial Resources (NIMR-2021-3).

Acknowledgments

The authors sincerely thank Prof. Aharon Oren (The Institute of Life Sciences, The Hebrew University of Jerusalem, The Edmond J. Safra Campus, 9190401 Jerusalem, Israel) and prof. Bernhard Schink (Department of Biology, University of Konstanz, D-78457 Konstanz, Germany) for their guidance in nomenclature.

Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Supplementary material

The Supplementary material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2022.1034816/full#supplementary-material

References

Aires, J. R., Köhler, T., Nikaido, H., and Plésiat, P. (1999). Involvement of an active efflux system in the natural resistance of Pseudomonas aeruginosa to aminoglycosides. Antimicrob. Agents Chemother. 43, 2624–2628. doi: 10.1128/aac.43.11.2624

Auch, A. F., von Jan, M., Klenk, H. P., and Göker, M. (2010). Digital DNA-DNA hybridization for microbial species delineation by means of genome-to-genome sequence comparison. Stand. Genomic Sci. 2, 117–134. doi: 10.4056/sigs.531120

Billi, D., and Potts, M. (2002). Life and death of dried prokaryotes. Res. Microbiol. 153, 7–12. doi: 10.1016/s0923-2508(01)01279-7

Blin, K., Shaw, S., Steinker, K., Villebro, R., Ziemert, N., Lee, S. Y., et al. (2019). antiSMASH 5.0: updates to the secondary metabolite genome mining pipeline. Nucleic Acids Res. 47, W81–W87. doi: 10.1093/nar/gkz310

Bric, J. M., Bostock, R. M., and Silverstone, S. E. (1991). Rapid in situ assay for indoleacetic acid production by bacteria immobilized on a nitrocellulose membrane. Appl. Environ. Microbiol. 57, 533–538. doi: 10.1128/aem.57.2.533-538.1991

Bianco, C., Imperlini, E., and Defer, R. (2009). Legumes like more IAA. Plant Signal. Behav. 4, 763–765. doi: 10.4161/psb.4.8.9166
Chaudhari, N. M., Gupta, V. K., and Dutta, C. (2016). BGPA: an ultra-fast pan-genome analysis pipeline. Sci. Rep. 6:24373. doi: 10.1038/srep24373

Collins, M. D., Pirooz, T., Goodfellow, M., and Minnikin, D. E. (1977). Distribution of menaquinones in actinomycetes and corynebacteria. J. Gen. Microbiol. 100, 221–230. doi: 10.1099/00221287-100-2-221

Dong, L., Ming, H., Yin, Y. R., Duan, Y. Y., Zhou, E. M., Nie, G. X., et al. (2014). *Rocconomas alkaliaterrace* sp. nov., isolated from an alkali geothermal soil sample in Tengchong, Yunnan, South-West China. Antonie Van Leeuwenhoek 105, 899–905. doi: 10.1007/s10482-014-1044-1

Dose, K., Bieger-Dose, A., Labusch, M., and Gill, M. (1992). Survival in extreme dryness and DNA-single-strand breaks. Adv. Space Res. 12, 221–229. doi: 10.1016/0273-1177(92)90176-x

Duca, D., Lory, J., Patten, C. L., Rose, D., and Glick, B. R. (2014). Indole-3-acetic acid in plant-microbe interactions. Antonie Van Leeuwenhoek 106, 85–125. doi: 10.1007/s10482-013-0095-y

Felsenstein, J. (1981). Evolutionary trees from DNA sequences: a maximum likelihood approach. J. Mol. Evol. 17, 368–376. doi: 10.1007/BF01733439

Felsenstein, J. (1985). Confidence limits on phylogenies: an approach using the bootstrap. Evolution 39, 783–791. doi: 10.1111/j.1558-5646.1985.tb00420.x

Foese1, B. U., Gossner, A. S., Drake, H. L., and Schramm, A. (2007). *Geminicoccus roseus* gen. nov., sp. nov., an aerobic phototrophic alphaproteobacterium isolated from a marine aquaculture biofilter. Syst. Appl. Microbiol. 30, 581–586. doi: 10.1016/j.syapm.2007.05.005

Gonzalez, C., Gutierrez, C., and Ramirez, C. (1978). Halobacterium vasuliortis sp. nov. an amylolytic and carbohydrate-metabolizing, extremely halophilic bacterium. Can. J. Microbiol. 24, 710–715. doi: 10.1139/m78-119

Kämpfer, P., Andersson, M. A., Jackel, U., and Salkinoja-Salonen, M. (2003). *Thiobacillus lupueritiae* gen. nov. sp. nov., and *Maricoccus roseus* gen. nov. sp. nov. representing two new genera of the alpha-1 subclass of the Proteobacteria. Syst. Appl. Microbiol. 26, 23–29. doi: 10.1016/S0176-177X(02)00035-8

Kim, O. S., Cho, Y. J., Lee, K., Yoon, S. H., Kim, M., Na, H., et al. (2012). Introducing ErTaxon-e: a prokaryotic 16S rRNA gene sequence database with phylogenies that represent uncultured species. Int. J. Syst. Evol. Microbiol. 62, 716–721. doi: 10.1099/ijs.0.038075-0

Kim, M., Oh, H. S., Park, S. C., and Chun, J. (2014). Towards a taxonomic coherence between average nucleotide identity and 16S rRNA gene sequence similarity for species demarcation of prokaryotes. Int. J. Syst. Evol. Microbiol. 64, 346–351. doi: 10.1099/ijs.0.059774-0

Kirmura, M. (1979). The neutral theory of molecular evolution. Sci. Am. 241, 98–100. doi: 10.1038/scientificamerican1179-98

Kirmura, M. (1980). A simple method for estimating evolutionary rates of base substitutions through comparative studies of nucleotide sequences. J. Mol. Evol. 16, 111–120. doi: 10.1007/BF01731581

Kluge, A. G., and Farris, J. S. (1969). Quantitative phyletics and the evolution of anurans. Syst. Zool. 18, 1–32. doi: 10.2307/2412407

Kroppenstedt, R. M. (1985). Fatty acid and menaquinone analysis of actinomycetes and related organisms. Soc. Appl. Bacteriol. Tech. Ser. 20, 173–199.

Li, W. J., Xu, P., Schumann, P., Zhang, Y. Q., Pakul, R., Xu, L. H., et al. (2007). *Georgonia ruanti* sp. nov., a novel actinobacterium isolated from forest soil in Yunnan (China), and emended description of the genus *Georgonia*. Int. J. Syst. Evol. Microbiol. 57, 1424–1428. doi: 10.1099/ijs.0.64749-0

Lowe, T. M., and Eddy, S. R. (1997). RNAscan-SE: a program for improved detection of transfer RNA genes in genomic sequence. Nucleic Acids Res. 25, 95–96. doi: 10.1093/nar/25.5.955

Martin, F. A., Posadas, D. M., Carrica, M. C., Cravero, S. L., O’Callaghan, D., and Zorreguieta, A. (2009). Interplay between two RND systems mediating antimicrobial resistance in *Brucella suis*. J. Bacteriol. 191, 2530–2540. doi: 10.1128/jb.00198-08

Meier-Kolthoff, J. P., Auch, A. F., Klenk, H. P., and Göker, M. (2013). Genome sequence-based species delimitation with confidence intervals and improved distance functions. BMC Bioinform. 14:60. doi: 10.1186/1471-2105-14-60

Minnikin, D. E., O’Donnell, A. G., Goodfellow, M., Alderson, G., Athalye, M., Schaal, A., et al. (1984). An integrated procedure for the extraction of bacterial isoprenoid quinones and polar lipids. *J. Microbiol. Methods* 2, 233–241. doi: 10.1016/0167-7012(84)90018-6

Nagakubo, S., Nishino, K., Hirata, T., and Yamaguchi, A. (2002). The putative response regulator BaeR stimulates multidrug resistance of *Escherichia coli* via a novel multidrug exporter system. *Mol. Microbiol.* 41, 461–467. doi: 10.1046/j.1365-2958.2002.02899.x

Reddy, G. S., Nagy, M., and Garcia-Pichel, F. (2006). *Belpnia moussenii* gen. nov., sp. nov., an alphaproteobacterium from biological soil crusts in the Colorado Plateau, USA. *Int. J. Syst. Evol. Microbiol.* 56, 51–58. doi: 10.1099/ijs.0.63764-0

Reinhold-Hurek, B., Hurek, T., Claeyssens, M., and van Montagu, M. (1993). Cloning, expression in *Escherichia coli*, and characterization of cellulosytic enzymes of *Azazara sp.*, a root-invasive diatostroph. *J. Bacteriol.* 175, 7056–7065. doi: 10.1128/JB.175.21.7056-7065.1993

Saitou, N., and Nei, M. (1987). The neighbor-joining method: a new method for reconstructing phylogenetic trees. *Mol. Biol. Evol.* 4, 406–425. doi: 10.1093/oxfordjournals.molbev.a040454

Shi, Y. L., Sun, Y., Jiang, Z. M., Ruan, Z. Y., Su, J., Yu, L. Y., et al. (2019). *Simplicispira lacundiana* sp. nov., a novel betaproteobacterium isolated from a freshwater reservoir. *Int. J. Syst. Evol. Microbiol.* 69, 129–133. doi: 10.1099/ijsem.0.035112

Tamura, K., Stecher, G., and Kumar, S. (2021). **MEGA11**: molecular evolutionary genetics analysis version 11. *Mol. Biol. Evol.* 38, 3022–3027. doi: 10.1093/molbev/msab20

Wang, Z., Fan, G., Hryc, C. F., Blaza, J. N., Serysheva, I. I., Schmid, M. F., et al. (2014). An allosteric transport mechanism for the AcrAB-TolC multidrug efflux pump. *elife* 4:e24905. doi: 10.7554/eLife.24905

Yoon, S. H., Ha, S. M., Lim, J., Kwon, S., and Chun, J. (2017). A large-scale evaluation of algorithms to calculate average nucleotide identity. *Antonie Van Leeuwenhoek* 110, 1281–1286. doi: 10.1007/s10482-017-8844-4