Diversity of Bacteria and the Characteristics of Actinobacteria Community Structure in Badain Jaran Desert and Tengger Desert of China

Ye Sun¹, Yun-Lei Shi¹, Hao Wang¹, Tao Zhang¹, Li-Yan Yu¹, Henry Sun² and Yu-Qin Zhang¹*

¹ Peking Union Medical College, Institute of Medicinal Biotechnology, Chinese Academy of Medical Sciences, Beijing, China, ² Desert Research Institute, Las Vegas, NV, United States

To assess the diversity of actinobacterial taxa in desert sands and obtain the novel microbial resources, 79 and 50 samples were collected from the Badain Jaran (BJD) and Tengger Deserts (TGD) of China, respectively. High-throughput sequencing (HTS) of environmental 16S rRNA genes within these samples was conducted on an Illumina MiSeq platform, using universal bacterial primers targeting the V3–V4 hypervariable region. Based on the HTS analyses, cultivation-dependent (CULD) techniques were optimized to identify the cultivable Actinobacteria members. A total of 346,766 16S rRNA gene reads comprising 3,365 operational taxonomic units (OTUs) were obtained from the BJD sands using HTS, while 170,583 reads comprising 1,250 OTUs were detected in the TGD sands. Taxonomic classification indicated that Actinobacteria was the predominant phylum, comprising 35.0 and 29.4% of the communities in BJD and TGD sands, respectively. Among the Actinobacteria, members of the Geodermatophilaceae were considerably abundant in both deserts, indicating that they represent ubiquitous populations within the deserts. At the genus level, Arthrobacter spp. and Kocuria spp. were dominant, and corresponded to 21.2 and 5.3% of the actinobacterial communities in BJD and TGD sands, respectively. Among the Actinobacteria, members of the Geodermatophilaceae were considerably abundant in both deserts, indicating that they represent ubiquitous populations within the deserts. At the genus level, Arthrobacter spp. and Kocuria spp. were dominant, and corresponded to 21.2 and 5.3% of the actinobacterial communities in BJD and TGD sands, respectively. A total of 786 and 376 actinobacterial strains were isolated and identified from BJD and TGD samples, respectively. The isolates comprised 73 genera of 30 families within the phylum Actinobacteria. In addition to the Geodermatophilaceae, Streptomyces spp. were a prominent component of the isolates, comprising 25% of the isolates from BJD and 17.5% of those from TGD. Comparison of the actinobacterial community structure in other ecosystems indicated that Geodermatophilaceae was the main actinobacterial group in desert sands, which is consistent with our results. Additionally, in these desert habits, Geodermatophilaceae and some other core groups may promote or inhabit the subsequent members’ occurrence or prosper to shape the bacteria community structure. However, it should be noted that a number of other low-abundance bacteria appear to be specific to desert sands, which
INTRODUCTION

Extremophiles have generated significant interest in the biological sciences in recent years, due to their unique genotypes, physiological functions, and secondary metabolites that hold great scientific and industrial value (Ciaramella, 1995). Deserts are extreme environments that are characterized by extreme aridity, intense solar UV radiation, and extreme shifts of temperature in day-night cycles. Consequently, deserts harbor numerous extremophiles (Subramani and Aalbersberg, 2013; Júlia et al., 2016). In addition to the above conditions, deserts feature chronic oligotrophy. Thus, sporadic vascular vegetation and microbiological crusts play critical roles in the primary production of this ecosystem, in addition to keeping soils from undergoing further desertification (Hawkes and Flechtner, 2002). Moreover, vegetation and microbiological crusts contribute greatly in shaping soil micro-ecosystem bacterial community structure in desert environments (Evans and Johansen, 1999; Belnap and Lange, 2003; Sun et al., 2015).

Deserts are typical of extreme, harsh ecosystems, where the availability of water is the cardinal parameter affecting organisms. Consequently, xerophilous microorganisms that are adapted to relatively high temperatures and radiation levels are likely to be the dominant populations in these ecosystems, including desiccation and radiation resistant phyla, such as Actinobacteria, Proteobacteria, and Bacteroidetes (Vikram et al., 2016).

The Badain Jaran Desert (BJD) is located in the northwest region of the Alxa Plateau of the Inner Mongolia Autonomous Region (39°20′–41°30′ N, 100°–104° E), and comprises an area of 49,000 km², making it the third largest desert in China (Yan et al., 2001). This region is known for its extreme continental climate and experiences the East Asian monsoon in its southwestern quadrant. The East Asian monsoon provides most of the precipitation for this area, accounting for 70% of the annual rainfall, which occurs within 10–35 days between July and September. The average annual rainfall is 150 mm while evaporation can reach as much as 1,500 mm. The Tengger Desert (TGD) is located in the southwest region of the Alashan Left Banner in the Inner Mongolia Autonomous Region and along the central border of Gansu Province (34°30′–39°00′ N, 103°–106°E). The TGD comprises a total area of about 43,000 km², and is the fourth largest desert in China. A typical continental climate dominates this region, with an annual average temperature of 7–9°C, and annual rainfall of 116–148 mm that primarily occurs in July and August. The annual evaporation of this region is 3,000–3,600 mm, and the average annual wind speed is 3–4 m/s. With increasing grassland degeneration and increases in desertification, the two deserts are almost contiguous with each other in some places (Figure 1).

Actinobacteria are a group of bacteria with high G+C content in the genomic DNA, and well-known for producing abundant secondary active metabolites. Since Waksman discovered streptomycin from Streptomyces sp. in 1943, actinobacterial members have been considered as an important source of new antibiotic-producing bacteria. Currently, about 70% antibiotics in clinical use are produced by various actinomycyces (Doul and Vining, 1990; Jose and Jebakumar, 2013). However, decades later, people found it is increasingly difficult to obtain novel compounds from the actinobacterial strains from normal environments. Therefore, more and more researchers proposed to find potential functional strains from various extreme environments, in which the external stress factors endow the microorganisms the unique defense mechanisms and metabolic systems, which may be more likely to produce novel antibiotics. Consequently, detecting various actinobacteria from extreme environments has become an important strategy for the discovery of new antibiotics (Phoebe et al., 2001; Wilson and Brimble, 2009). Till now, no systematic studies on actinobacteria regarding their potential abilities to produce bio-active substances and their ecological function in deserts have been reported.

The primary goal of the present research was to obtain a systematic understanding of the bacterial community structure and diversity of actinobacteria inhabiting the two immense BJD and TGD deserts of China. Further, we attempted to better characterize the distribution characteristics of actinobacterial population in these environments. Based on the above findings, we intended to optimize the isolation and cultivation strategies to explore the specific actinobacterial strains and to assess their bio-characteristics, potential functions and applications in depth.

MATERIALS AND METHODS

Sample Collection

A total of 79 samples were collected from the BJD (38°20′–41°30′N, 100°–104°E). Sporadic vascular vegetation and small salt lakes characterize this desert. We thus assigned samples to the following three types based on vegetation cover and salt lake presence: sands with sporadic plants (SP), sands without any vegetation (NV), and sands around salt lakes (SL). Detailed information of samples including types, and the specific collection locations are provided in Table S1. Fifty samples were collected from the TGD, where the microbial...
crusts are generally categorized into the following three types: Cyanobacteria-dominated crusts (CC), moss-dominated crusts (MC), and lichen-dominated crusts (LC). Samples from the TGD also included bare sands (BS) (Sun et al., 2015), and sample information is provided in Table S2.

All samples were placed in sterilized envelopes following collection and then taken to the laboratory within 3 days of collection. The samples were immediately processed for cultivation assays and total DNA extraction after arriving at the laboratory.

Isolation and Cultivation Media
Isolation Media
The following seven media were prepared to isolate microbial strains:

M1: 1/5 strength R2A (Difco).
M2: 5 g/L yeast extract, 2 g/L cellobiose, 2 g/L CaCO₃, 0.5 g/L MgSO₄·7H₂O, 1 g/L K₂HPO₄, and 15 g/L agar.
M3: 0.1 g/L NH₄NO₃, 2 g/L sodium propionate, 0.05 g/L MgSO₄·7H₂O, 0.1 g/L KCl, and 15 g/L agar.
M4: 1 g/L humic acid, 1 g/L asparagine, 0.01 g/L FeSO₄, 0.5 g/L Na₂HPO₄, 1.7 g/L KCl, 0.02 g/L CaCO₃, and 15 g/L agar.
M5: 2 g/L trehalose, 5 g/L yeast extract, 2 g/L CaCO₃, 0.5 g/L MgSO₄·7H₂O, 1 g/L K₂HPO₄, and 15 g/L agar.
M6: 0.5 g/L K₂HPO₄, 0.25 g/L yeast extract, and 15 g/L agar.
M7: 5 g/L yeast extract, 3 g/L peptone, 10 g/L glycerol, 1.25 g/L sodium pyruvate, 1.25 g/L glycine betaine, and 15 g/L agar.

Media were adjusted to pH 7.2–7.5 using 1 M HCl and/or 1 M NaOH. In addition, betaine (0.125% w/v), sodium pyruvate (0.125% w/v), compound trace salts solution (0.1% w/v), and compound vitamins (0.1% w/v) were added to the media as described in Sun et al. (2015). Nystatin (25 mg/L) and potassium dichromate (50 mg/L) were added to the media to inhibit the growth of fungi and Gram negative bacteria.

Cultivation Media
Based the observation of the colony diversity growth in the isolation media, the PYG medium was selected to cultivate isolates after initial isolation and consisted of 3 g/L peptone, 10 g/L glycerol, 5 g/L yeast extract, 1.25 g/L sodium pyruvate, 1.25 g/L glycine betaine, and 15 g/L agar, adjusted to pH 7.5. PYG medium was supplemented with compound trace salts solution (0.1% v/v), and compound vitamins (0.1% w/v).

Total DNA Preparation From Sand Samples and PCR Amplification
The 79 sand samples from BJD were pooled into seven composite samples, and the 50 sand samples from TGD were pooled into eight composite samples, according to the environments where the sands were collected. Detailed information of the 15 samples is provided in Table S3. Total genomic DNA from each of the 15 pooled samples was extracted with a PowerSoil DNA isolation kit (MoBio, USA) according to the manufacturer’s protocols. Total DNA was then used as template for PCR amplification of 16S rRNA genes. The V3 to V4 hypervariable regions of 16S rRNA genes were PCR amplified using the universal bacterial primers 5′-ACTCCTACGGGAGGCAGCAG-3′ (338F) and 5′-GGACTACHVGGGTWTCTAAT-3′ (806R). PCR amplifications were performed using high fidelity TransStart Fastpfu DNA Polymerase (Transgen, China) in 20 µL reaction mixtures containing 4 µL of 5× FastPfu Buffer, 2 µL of 2.5 mM dNTPs, 0.8 µL of each primer (5 µM), 0.4 µL of FastPfu Polymerase, and 10 ng of template DNA. The reaction cycling conditions consisted of the following steps: 5 min of an initial denaturation at 94°C
followed by 35 cycles of denaturation at 94°C for 30 s, 45 s of primer annealing at 55°C, 40 s of elongation at 72°C, and then a final 10 min elongation at 72°C.

**Illumina MiSeq Sequencing and Raw Data Preprocessing**

An AxyPrep DNA Gel Extraction Kit (Axygen, USA) and Quantifluor™-ST system (Promega, USA) were used to purify and quantify amplicons, respectively. Purified amplicons were pooled in equimolar concentrations and sequenced on an Illumina MiSeq platform with paired-end sequencing (2 × 250 bp). The QIIME software package (version 1.18; White et al., 2009; Caporaso et al., 2010) was used to demultiplex and quality-filter raw reads by removing reads meeting the following criteria: (a) 300 bp reads exhibiting any 50-bp sliding window with an average quality score < 20, and discarding truncated reads that were shorter than 50 bp; (b) any barcode mismatch, greater than two nucleotide mismatches in primers, and reads containing ambiguous characters (c). Lastly, paired sequences were assembled based on their overlapping sequences that were longer than 10 bp. Unassembled reads were discarded.

**Statistical Analyses**

Chimeras were identified and removed using the UCHIME program (Robert et al., 2001). The Unite database (Release 7.0, http://unite.ut.ee/index.php; Köljalg et al., 2013) was used as the taxonomic reference database to assess the taxonomic affiliation of each 16S rRNA gene sequence using a confidence threshold of 70% and the RDP Classifier (Wang et al., 2007). Sequences were clustered into operational taxonomic units (OTUs) using a 97% similarity threshold in the UPARSE program (version 7.1, http://drive5.com/uparse/). Alpha diversity and beta diversity metrics were calculated based on OTU abundances within and among samples. Specifically, alpha diversity was measured using the Chao1 richness estimator, and Shannon's diversity index, while measuring the coverage of richness using Good's coverage. Principal co-ordinates analysis (PCoA) was used to assess differences in OTU composition among samples. An analysis of similarities (ANOSIM) test was conducted to determine if sample types contained significantly different bacterial communities. Metastats and LEfSe (LDA effect size) analysis were employed to identify the bacterial groups that were significantly differentiated among sample types. Lastly, co-occurrence networks were constructed to visualize the OTU-based similarities among different communities (White et al., 2009; Caporaso et al., 2010).

**Strains Isolation, Purification, Maintenance, and Identification**

The dilution plating method was used to isolate microbial strains from the sand samples. Approximately 0.3 mL of a 10⁻⁴ dilution of a sand sample suspension was spread on each isolation plate. Every kind of media was parallelly spread with the suspension on two same plates and then incubated at 28°C and 45°C for 21 days, respectively. Single colonies were transferred to freshly prepared PYG plates that were supplemented with compound trace salts solution and compound vitamins, followed by subsequent purification of isolates. Purified isolates were maintained on PYG slants at 4°C and also in glycerol suspensions (20%, v/v) at −80°C (Yue et al., 2006). Genomic DNA was extracted from pure cultures and PCR amplification of 16S rRNA genes was conducted as described in Xu et al. (2003). The sequences of the isolates were compared with available 16S rRNA gene sequences from GenBank using the BLAST program and the EzBioCloud (https://www.ezbiocloud.net/) (Yoon et al., 2017).

**Antimicrobial Activity Screening**

Antimicrobial activities of the isolates were investigated by using media containing *Escherichia coli* ATCC 25922, *Pseudomonas aeruginosa* ATCC 27853, *Enterococcus faecalis* ATCC 29212, *Klebsiella pneumoniae* subsp. *pneumoniae* ATCC 700603, and *Candida albicans* ATCC 10231, with a concentration of 10⁶ colony forming units (CFU) per mL. These assays were performed using Kirby Bauer Method with a culture broth concentration of 1% (v/v).

**RESULTS**

**Sequences and Data Information**

The SRA accession number in DDBJ/EMBL/GenBank for the sequences data is SRP134260.

**Bacterial Richness and Diversity**

A total of 349,374 reads with an average of 23,291 reads per sample remained after quality filtering, and were clustered into 4,298 OTUs at the 97% sequence similarity level. The alpha diversity of each sample estimated by the Chaol estimator, and Shannon's index, in addition to the coverage estimated by Good's coverage, are provided in Table 1. Rarefaction analyses using the Shannon index as a diversity metric indicated that our sequencing efforts covered nearly all of the diversity that would be expected to be found in these samples (Figure S1). On average, 923 (95% CI: 799–1,029) and 900 (95% CI: 653–1,165) OTUs were found in each of the BJD and TGD samples, respectively. Samples from the desert sands exhibited almost identical Chao1 and Shannon index values, as indicated by ANOVA tests, suggesting no significant difference in the richness or diversity of bacteria between the two deserts (Figure 2).

Among the BJD samples, SP samples had the highest Chao1 values, whereas samples from NV, SL and SP did not significantly differ. Among the TGD samples, bacterial richness and diversity was slightly higher in BS samples compared to the others, which was reflected in higher Chaol and Shannon index values (Figure 3).

**Bacterial and Actinobacterial Community Structure**

A total of 32 phyla, 97 orders, and 470 genera were identified in BJD samples, while 28 phyla, 73 orders, and 302 genera were identified in TGD samples. *Actinobacteria* dominated the bacterial communities of both deserts (35 and 29% relative abundance in BJD and TGD, respectively), followed by *Proteobacteria, Bacteroidetes*, and *Chloroflexi* (Figure 4). *Frankiales, Micrococcaceae, Micromonosporales,* and...
Acidimicrobiales were the most abundant orders within the 15 sand samples. Among these, the families Geodermatophilaceae, Micrococccaceae, Micromonosporaceae, and Rubrobacteriaceae were the predominant actinobacterial populations in both deserts. Lastly, the most abundant genera were Arthrobacter and Rubrobacter in the actinobacterial community, followed by Blastococcus, Modestobacter, and Geodermatophilus, which are members of the Geodermatophilaceae (Figure 5). In BJD, Micrococccaceae was much more prevalent than Geodermatophilaceae, while the reverse was observed in TGD samples. Arthrobacter was the most abundant genus in BJD samples and the second most abundant genus in TGD samples. Kocuria was a dominant genus in BJD samples, but only a few Kocuria OTUs were detected in TGD samples. Conversely, Actinoplanes were more abundant in TGD samples than in BJD samples. In addition to these dominant genera, a large number of rare bacterial genera were also detected (we defined rare bacterial groups as less than 0.1% abundance of communities), and included the genera Conexibacter, Longispora, Dactylosporangium, Umezawae, Demequina, Janibacter, and Motilibacter in BJD and genera Ornithinimicrobium, Angustibacter, Nakamuraella, Aquiluna, Williamsia, Amycolatopsis, and Kineospora in TGD (Figure 5).

Metastats analysis indicated that the composition of actinobacterial genera in the two deserts were not significantly different (P = 1) (Table S4). However, different sand types represented distinct ecosystems, and generally, bacterial community composition was more strongly correlated with sampling site rather than micro-ecosystem type (Figure 6).

In particular, ANOSIM (R = 0.328, p = 0.001) indicated that community structures were not different between samples taken from sampling sites that were close together. LEfSe analyses indicated that there were some significant differences in community composition among the three sample types from BJD. For example, the abundances of the genera Virgisorangium, Nocardiopsis, Brachybacterium, Kineococcus, Georgenia, and Streptosporangium were highly differential among sample types within BJD (Figure 7). However, no significant differences in the abundances of these genera were observed for samples from the TGD. The Geodermatophilaceae and Micrococccaceae were the dominant families in NV, SL, SP, and BS samples, whereas Micromonosporaceae were more abundant in the LC, CC, and MC samples. Further, Pseudonocardia were more abundant in NV, SP, BS, LC, CC, and MC samples, but not in SL samples. Arthrobacter was the predominant genus in NV, SL, SP, and BS samples, but not in LC, CC, or MC samples. Lastly Rubrobacter were abundant in LC, CC and MC samples, but not NV, SL, SP, or BS samples (Figure S2).

We used co-occurrence network analysis to identify correlations between different genera among samples. A total of 27 nodes and 84 correlation edges were used to represent the dominant genera (>1% relative abundance). Correlation analysis indicated 77 positive and seven negative correlations. The genera Solirubrobacter and Actinoplanes exhibited the most correlations (12 edges for each), followed by the Pseudonocardia, Marmoricola, Saccharothrix, and Friedmanniella which each comprised 10 edges. These results indicated that the six genera were likely to be the key actinobacterial members in the sand communities of the two deserts. Among the family Geodermatophilaceae, three genera appeared to also be important with Geodermatophilus exhibiting six edges, Modestobacter with five, and Blastococcus with two (Figure 8).

### Isolation of Actinobacterial Strains
A total of 786 actinobacterial strains were isolated and identified from the BJD samples. 16S rRNA gene sequencing indicated that these 786 strains belonged to 30 families and 73 genera (Figure S3A). In addition, 376 actinobacterial strains were isolated from TGD samples, comprising 18 families and 29 genera (Figure S3B). 16S rRNA gene sequencing indicated that members of Geodermatophilaceae were the major cultivated actinobacterial taxa in both of these deserts, which is consistent with the cultivation-independent analyses. Moreover, the taxonomic distributions of isolates were highly similar to those of the cultivation-independent analyses.

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**Table 1** Summary of high-throughput sequencing data from the 15 samples.

| Sample ID | Sample type | Number of OTUs | Good's estimator (%) | Chao1 | Shannon |
|-----------|-------------|----------------|----------------------|-------|---------|
| BD201610S1 NV | 799 | 98.65 | 1,046 | 4.95 |
| BD201610S2 NV | 938 | 98.51 | 1,145 | 5.3 |
| BD201610S3 NV | 925 | 98.08 | 1,190 | 5.22 |
| BD201610S4 SL | 804 | 99.31 | 1,051 | 5 |
| BD201610S5 SL | 798 | 99.19 | 1,172 | 4.74 |
| BD201610S6 SP | 1,129 | 99.49 | 1,294 | 5.6 |
| BD201610S7 SP | 1,067 | 99.29 | 1,283 | 4.91 |
| SPT8001BS BS | 1,155 | 98.72 | 1,401 | 5.42 |
| SPT8002BS BS | 1,165 | 98.72 | 1,405 | 5.67 |
| SPT8003LC LC | 719 | 98.66 | 998 | 4.95 |
| SPT8004LC LC | 824 | 98.89 | 1,015 | 4.34 |
| SPT8005CC CC | 1,039 | 98.58 | 1,463 | 5.57 |
| SPT8006CC CC | 863 | 99.34 | 852 | 4.32 |
| SPT8007MC MC | 892 | 98.57 | 1,108 | 5.34 |
| SPT8008MC MC | 756 | 98.98 | 1,063 | 3.42 |

BS, Bare sand; CC, Cyanobacteria-dominated crusts; MC, Moss-dominated crusts; LC, Lichen-dominated crusts; SP, sand soil with sporadic plants; NV, sands without any vegetation; SL, sand sample from around salt lakes.
Arthrobacter spp. and Kocuria spp. were the primary cultivated genera, and comprised 13.4 and 4.1% of the actinobacterial community isolates in BJD, respectively (Figure S4; Figure 9).

Streptomyces spp. were also dominant among the isolated genera, comprising 25% of BJD isolates and 17.5% of TGD isolates. In addition, some low abundance taxa that were
detected using cultivation-independent analysis including *Deinococcus* spp., *Skermanella* spp., *Actinophytocola* spp., *Patulibacter* spp., *Rufibacter* spp., and *Geminicoccus* spp. were also isolated.

**Comparative Analysis of HTS and CULD Results**

Of the dominant genera (>1%), 4.3 and 32.2% were detected using the two methodologies in BJD and TGD samples, respectively (*Figure 10*). In addition, 14.0 and 17.1% of the rare genera (<0.1%) were identified by both methodologies in BJD and TGD, respectively. However, there were some genera that were only detected by either cultivation-dependent (CULD) or high-throughput sequencing (HTS) analysis (*Figure 10*). These results indicate that the composition of the native microbial communities are not only driven by the dominant groups, but also rare taxa that are active and integral members of community structure.

**Actinobacterial Members in Desert Sands**

*Geodermatophilaceae* populations were the dominant group in both the BJD and TGD deserts, comprising 27.4 and 18.2% of the *Actinobacteria* according to cultivation-independent analyses. The genera *Blastococcus*, *Modestobacter*, and *Geodermatophilus* exhibited nearly equivalent diversity in the two deserts (*Figure S4*). However, *Blastococcus* was more ubiquitous and
abundant in all samples, compared to the other two genera. *Modestobacter* was more abundant in NV, MC, and CC samples, while OTUs associated with this genus were rare in SL, SP, and BS samples. *Geodermatophilus* was abundant in NV, SP, and CC samples, but not in LC or MC samples. Generally, NV, SP, BS, and CC samples harbored the most abundant *Geodermatophilaceae* communities (Figure 11). Further, more *Geodermatophilaceae* isolates were obtained from the NV and BS samples, which could be regarded as more oligotrophic environments. In total, 52 *Geodermatophilaceae* strains were obtained from BJD, including 36 *Blastococcus* spp., six *Geodermatophilus* spp. and 10 *Modestobacter* spp. In addition, 34 *Blastococcus* spp., 11 *Geodermatophilus* spp., and 25 *Modestobacter* spp. were obtained from TGD. Importantly, 21 actinobacterial isolates from BJD representing novel species within the following 15 genera were obtained: *Actinokineospora*, *Actinomycetospora*, *Auraticoccus*, *Blastococcus*, *Cellulomonas*, *Kibdelosporangium*, *Mariniluteicoccus*, *Microbacterium*, *Nesterenkonia*, *Nocardia*, *Nocardiooides*, *Pseudonocardia*, and *Solirubrobacter*. Likewise, 16 novel taxa were isolated from TGD that were distributed among 11 genera (Figure 12). These novel actinobacterial taxa are worthy of further bioprospecting studies.

**Antimicrobial Activities**

Five-hundred actinobacterial isolates from BJD were screened for antimicrobial activities. Of these, 1.6% of the isolates exhibited antimicrobial activity against *Escherichia coli* ATCC 25922, 3.5% against *Enterococcus faecalis* ATCC 29212, 1.7% against *Klebsiella pneumonia* subsp. *pneumoniae* ATCC 700603, 1.2% against *Pseudomonas aeruginosa* ATCC 27853, and 2.2% against *Candida albicans* ATCC 10231. Overall, 10.36% of the tested isolates from BJD exhibited antimicrobial activity in one or more screens.

**DISCUSSION**

With the rapid technological innovation in molecular biology in recent years, various new techniques, and in particular high-throughput cultivation-independent approaches, have been extensively applied in microbial ecology research. These new methodologies have greatly expanded our knowledge of bacterial community composition. Consequently, numerous studies have shown that microbial diversity is far higher and more complex than previously thought. 16S rRNA gene cloning and denaturing gradient gel electrophoresis (DGGE) have been traditionally used to identify uncultured microbial community structures in the past. However, these methods are particularly susceptible to 16S rRNA gene copy numbers, and the methods are both complicated and costly. High-throughput sequencing precludes the need to build clone libraries, and the efficiency associated with HTS sequencing approaches and the widespread adoption of the Illumina MiSeq sequencing platform has accelerated analytical capabilities. Consequently, HTS analysis on the Illumina MiSeq platform has provided comprehensive and efficient analyses of environmental microbial communities. (Navarro-Noya et al., 2013; Poulsen et al., 2013; Sul et al., 2013; Yu et al., 2013; Wu et al., 2014). Claesson et al. (2010) conducted a comprehensive
analysis of 9 variable regions of the 16S rRNA gene. The results showed that V1 and V9 had the worst effects, followed by V7 and V8 regions, while V3, V4, and V5 regions were relatively better. In addition, by comparing the regions of V1/V2, V2/V3, V3/V4, V5/V6, and V7/V8 in series, the accuracy of the tandem region V3/V4 and V4/V5 were the highest. Considering the classification efficiency (CE), the V3 / V4 region had significant amplification preference relative to other regions (Claesson et al., 2010). Here, we analyzed the bacterial communities of two Chinese deserts using HTS, and accordingly, modified the
media and temperature for isolation and cultivation the targeted actinobacterial strains. Our results provide evidence for a number of actinobacterial strains that are specific for these desert sands and were isolated and identified via cultivation-dependent and -independent analyses. Importantly, these results describe the diversity and composition of actinobacterial communities in different sand types, which has not been systematically investigated in these two deserts (Sun et al., 2015; Li et al., 2016).

Comparative results from the two deserts and the different mico-ecosystems indicated that geographical barrier contributed much more in shaping the bacterial communities than micro-ecological types. On the other side, even in a certain same terrain, the indigenous microorganisms’ ecological service function resulted in discriminating micro-ecosystem.

Deserts are typical of extreme, harsh ecosystems, where the availability of water is the cardinal parameter affecting organisms. Consequently, xerophilous microorganisms that are adapted to relatively high temperatures and radiation levels are likely to be the dominant populations in these ecosystems. Actinobacteria are a ubiquitously distributed bacterial phylum inhabiting diverse terrestrial ecosystems. Based on this observation, we approved that Actinobacteria is the predominant microorganisms in desert sand microbial communities using HTS and CULD data. Generally, Actinobacteria, characterized by high genomic GC content, complex peptidoglycan layer containing in the thick cell wall, motility with spore-bearing cells and pigmentation in colonies, may have evolved into some genetic and metabolic functions to help them inhabit in the deserts with drought and high ultraviolet radiation (Mohammadipanah and Wink, 2012).

A total of 4,298 OTUs were obtained using cultivation-independent HTS, indicating a high level of bacterial diversity in these desert environments. Importantly, the cultivation-independent and -dependent analyses provide evidence for valuable microbial resources that warrant further research.

FIGURE 9 | Pie chart showing the percentage of relative abundance of bacterial communities from two desert sand samples in CULD method. (A) Badain Jaran desert sands; (B) Tengger desert sands. The sequences of the bacteria were assigned to family (1/3) and genus level (2/4).
In particular, some specific, novel Actinobacteria members and rare microbial groups were identified that could serve as highly valuable microbial resources. OTUs affiliated with the Geodermatophilaceae were frequently detected among the 15 sample types analyzed here, which is consistent with our previous supposition that Geodermatophilaceae are ubiquitous within arid deserts (Sun et al., 2015). Geodermatophilaceae have been detected from various relatively extreme environments, including stone habitats (Salazar et al., 2006; Chouaia et al., 2012; Gtari et al., 2012; Normand et al., 2012), dry hot valleys (Nie et al., 2012), and arid sand from deserts (Montero-Calasanz et al., 2012; Montero-Calasanz C. et al., 2013; Montero-Calasanz M. C. et al., 2013; Saul-Tcherkas et al., 2013), among others. Thus, Geodermatophilaceae strains may be pioneer microorganisms that play a key role in shaping the bacterial community structure in these harsh environments. Furthermore, to some extent, in these desert habits, the microbial community structures may always be doomed from some pioneer population’s colonization, where the core groups may promote or inhabit the subsequent members’ occurrence and prosper.

Actinoplanes and Kocuria were abundant members of their communities and were also two key genera of the desert sand communities based on co-occurrence network analyses. Interestingly, Actinoplanes and Kocuria were negatively correlated in the co-occurrence network. A large number of
FIGURE 12 | Dendrogram based on 16S rRNA gene sequences analysis of 37 novel species and some relevant strains in the two deserts. Bar, 100 substitutions per nucleotide position.
Kocuria spp. were isolated from our cultivation assays (21 strains) from sand samples, yet only a few Actinoplanes strains were obtained. In addition, Streptomyces was an abundant component of both the HTS and cultivation assays. However, Streptomyces and Actinoplanes were negatively correlated in the co-occurrence network analysis. These results point to a potentially intriguing negative interaction between Streptomyces, Kocuria, and Actinoplanes that may explain the paucity of Actinoplanes isolates that were obtained. Future research should investigate the potential for supplements that inhibit the growth of Streptomyces and Kocuria in order to promote the isolation of Actinoplanes. Regardless, our results suggest the presence of valuable microbial strains, particularly rare community members that are present within these desert sands, and which may hold great significance in understanding the ecology of these systems or in novel microbiological drug research. The analyses here, particularly the co-occurrence networks, provide a framework from which to design isolation media that may promote the growth of these potentially valuable taxa.

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AUTHOR CONTRIBUTIONS

YS carried out the experiments and prepared the manuscript. Y-LS helped prepare some experiments. TZ, L-YY, and HW contributed in sampling from deserts. HS helped correct the writing. Y-QZ is responsible for designing the research and preparing the manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicro.2018.01068/full#supplementary-material

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