Sand Dune *Streptomyces* JB66 Native to the Great Indian Thar Desert Inhibits Multidrug-Resistant Pathogens

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ABSTRACT

The efficacy of an actinomycete strain JB66 recovered from a sand dune soil from the Bikaner district of the Thar desert in inhibiting the growth of various bacterial pathogens was studied. The type strains *Staphylococcus aureus*, *Shigella flexneri*, *Klebsiella pneumoniae* and the clinical isolates *Escherichia coli*, multidrug-resistant *S. aureus* and *P. vulgaris* were included in the antimicrobial assays. Polyphasic characterization of JB66 isolate revealed its identity as *Streptomyces* (MH762010). It showed 88.99-89.24% sequence similarity with the other members of this genus and share the maximum (88.89%) similarity with *Streptomyces* sp. ATSC13. The strain JB66 was found to produce a high amount of extracellular L-asparaginase, catalase, gelatinase, protease, tyrosinase and urease enzymes. The partial chemical categorization of the methanolic crude extract of the JB66 strain led to the preliminary identification of various metabolic compounds. The thin-layer chromatography fractionation revealed the presence of prodigiosin pigment or chandramycin, cephalosporin or zeatin, daidzein, demethoxy rapamycin, 4,6-dihydroxy-7-methoxyisoflavone, munumbicins and amiclenomycin like compounds. Bio-autography revealed that the metabolites localized at the Rf values of 0.40, 0.46 and 0.53 in TLC profile had the actual bioactive fractions. UV-VIS spectrum absorbance maxima at 288 nm revealed the presence of an aromatic nucleus.

Keywords: Actinomycetes, Thar desert, Sand dune, MDR, TLC, Rapamycin, Cephalosporin.

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INTRODUCTION

Nature is a prolific source of many useful natural products that play many prominent roles in the development of new therapeutic agents. [1-3] A huge proportion of structurally different bioactive metabolites are synthesized by microorganisms to sustain in different ecosystems. [1, 4] Within microorganisms, actinobacteria represent the most promising group of bacteria that synthesize approximately two-thirds of all known natural bioactive products including some of the compounds most widely used as anticancer, anti-inflammatory, antibacterial, antifungal and antitumour agents etc. [5-16] Indeed, an estimated 70% or maybe more antibiotics...
currently used at the commercial levels are produced by the members of the genus *Streptomyces* in the phylum actinobacteria.\(^5,7-10\) Due to the emergence of multidrug-resistant pathogens against existing antibiotics, it is highly important to find the novel source of antibiotics and alternate methods on an urgent basis. By exploring the unexplored or less explored environment, the possibility of finding new biologically active metabolites can be increased.\(^19-21\) In this context, the less explored desert environment might be considered useful for finding novel actinobacterial bioactive metabolites.\(^22-26\) Therefore, in the present study, sampling was done from a sand dune soil in the Bikaner district of the Thar Desert for the isolation and identification of actinobacteria producing useful antibiotics against multidrug-resistant pathogens. Intriguingly, based on the antagonistic activity against multidrug-resistant pathogens, a *Streptomyces* strain JB66 was identified producing cephalosporin and rapamycin like compounds.

**MATERIAL AND METHODS**

**Soil sampling and isolation of actinobacteria**

The samples were collected from a sand dune soil from the Bikaner region of the Thar desert in Rajasthan. The samples were collected at the depth of 6-10 cm using a sterile scoop in a sterile zipped bag. The collected samples were transferred to the laboratory for further analysis. To avoid the growth of undesired bacteria and for the selective isolation of actinomycetes, all samples were subjected to various pretreatment procedures before serial dilution. Pretreatment methods, drying\(^27-28\) and moist heating\(^29\) were employed. A pretreated soil sample was added to 50 mL of sterile distilled water in a 250 mL Erlenmeyer flask. This was kept for shaking on an orbital shaker for 30 min at 37°C and serial dilutions were made. From each dilution 0.1 mL of sample was spread on different selective isolation media namely ISP2 (International Streptomycete Project),\(^30\) Benedict’s modification of the Lindebein\(^31\), Czapek’s\(^32\), Gauze’s\(^33\), humic-acid vitamin\(^34-35\) potassium-tellurite\(^36\), raffinose-histidine\(^37\), R2YE\(^38\), starch-casein\(^39\) and MAS\(^18\) agar media. The antibiotics kanamycin, nalidixic acid and cyclohexamide (25µg/mL) were added in the media to inhibit bacterial and fungal contaminations, respectively. The inoculated plates were incubated at 37°C for 7-20 days. Following incubation, the appearance and growth of microorganisms were observed every day. After isolation, the isolates were sub-cultured to ensure their purity and maintained on MAS agar medium and preserved using 20% (v/v) glycerol at -40°C.\(^33\)

**Screening for antibacterial activity**

The putative actinomycete isolates were screened for their antibacterial activity using the agar plug method\(^40\) in primary screening. The type strains *Staphylococcus aureus* (NCIM 2079), *Shigella flexneri* (NCIM 4924) and *Klebsiella pneumoniae* (NCIM 2719) and one clinical isolate *Escherichia coli* (clinical isolate responsible for Urinary Tract Infection (UTI)) were used as the pathogens in antimicrobial assays. A fully grown culture of a pathogenic strain was seeded by swabbing on the MAS agar medium. After that, agar plugs were cut from 10 days old actinomycete agar culture and placed on the agar plate seeded with different pathogens in the separate experiments. The plates were incubated overnight at 37°C for 24-72 h. Inhibition zones around the plugs after the incubation period were observed. The agar well diffusion method was used in the secondary screening to assess the antimicrobial activity of putative actinobacterial isolates. The isolates were grown in MAS broth medium. After centrifugation at 10,000 g for 10 min, the culture supernatant was introduced in a well bored on the agar plate seeded with different pathogens. Consequently, the inhibitory zone produced was measured. During the time course of fermentation, antibiotic production, antimicrobial activity at different time points were also monitored. The isolate JB66 that exhibited broad-spectrum antimicrobial activity against the tested pathogens was selected for further studies.

**Fermentation and biological activity of JB66 against MDR pathogens**

The actinomycte isolate JB66 exhibiting the highest antagonistic activity against all the tested pathogen was cultured in ISP2 medium and incubated in a rotatory shaker at 120 rpm at 30°C for 10 days. Subsequently, the culture broth was centrifuged (6000 g, 15 min) and the supernatant was extracted with an equal volume of ethyl acetate (EA) at 1:1 (v/v) ratio. The extract was concentrated to dryness at 50°C. Methanol was used to dissolve the crude extract and stored at 4°C until further use. The biological activity of the crude methanolic extract of actinobacterial isolate JB66 was tested against multidrug-resistant Gram-positive *S. aureus* and Gram-negative *P. vulgaris* (clinical isolates). Antibiotic-resistance profile of *S. aureus* was determined against amoxicillin-clavulanic acid (30µg), ampicillin (10µg), azithromycin (30µg), cefepime (30µg), ceftiraxone (30µg), cefotaxime (30µg), cefuroxime (30µg), cephalxin (30µg), ciprofloxacin (1µg), clindamycin (2µg), cloxacillin (30µg), trimethoprim-sulfamethoxazole (25µg), erythromycin (15µg), imipenem (10µg), levofloxacin (5µg), linezolid (30µg), meropenem (10µg), piperacillin-tazobactam (100/10µg), ticoplacin (30µg), tetracycline (30µg), and vancomycin (30µg). Whereas antibiotic-resistance profile of *P. vulgaris* was determined against amikacin (30µg), amoxicillin-clavulanic acid (30µg), aztreonam (30µg), cefepime (30µg), Cefoperazone-Sulbactam (75/30µg), ceftiraxone (30µg), cefotaxime (30µg), cefuroxime (30µg), ciprofloxacin (1µg), trimethoprim-sulfamethoxazole (25µg), ertapenem (10µg), gatifloxacin (5µg), gentamicin (10µg), imipenem (10µg), levofloxacin (5µg), meropenem (10µg), netilmicin (30µg), norfloxacin (10µg), ofloxacin (5µg), piperacillin-
tazobactam (100/10μg). The interpretation of their antibiotic susceptibilities was made using CLSI guidelines (M100-S22). [41]

**Scanning electron microscopy**

After the assessment of antimicrobial activity, the isolate JB66 that exhibited the highest anti-microbial activity against MDR pathogens was further subjected to its external morphological studies using scanning electron microscopy. The sample was prepared and analyzed according to the protocol described. [42]

**Phenotypic and biochemical characterization**

Morphological characterization of presumptive actinomycete isolate JB66 was performed by growing it on the ISP media as described by Shirling and Gottlieb (1966). [30] The growth characteristics of 7 to 14 days old JB66 culture were determined using yeast extract/malt extract agar (ISP2), oatmeal agar (ISP3), inorganic salts/starch agar (ISP4), glycerol-asparagine agar (ISP5), peptone-yeast extract iron agar (ISP6) and tyrosine agar (ISP7) media and the colour of aerial and substrate mycelia was noted. The colonial morphology of the strain was examined by slide culture method. [33]

Physiological characterization at variable pH range (5-11), temperature (25-55°C) and salinity tolerance at different NaCl strength (0-10%) was done. Briefly, in different assays, the isolate JB66 was inoculated into the broth tubes containing different salt concentrations and variable pH and incubated at 45°C for 7 days. The pH of the MAS broth medium was adjusted with 0.1N NaOH/0.1N HCl. For temperature tolerance, the broth medium inoculated for JB66 strain was incubated at 25°C, 35°C, 45°C and 55°C separately. Carbohydrate utilization was determined by growing the isolate in ISP-9 medium [43] by using 1% of various carbon sources viz., arabinose, dextrose, fructose, galactose, D-glucose, lactose, mannitol, mannose, meso-inositol, pectin, rhamnose, ribose, salicin, sucrose, starch and D-xylene. The utilization of various nitrogen sources (1%) was also studied by using alanine, aspartic acid, arginine, asparagine, cysteine, glutamic acid, glycine, histidine, isoleucine, methionine, proline, tyrosine, tryptophane in the basal medium consisted of (g/L) K2HPO4 1.0 g, MgSO4.7H2O 0.5 g, CaCl2.2H2O 0.04 g, FeSO4.7H2O 0.005 g, ZnSO4.7H2O 0.0005 g, 15 g agar and 1.0% of each of the nitrogen sources. The strain JB66 was screened for the production of extracellular catalase, amyylase, cellulase, gelatinase, L-asparaginase, lipase, tyrosinase and urease enzymes.

**Molecular characterization**

Identification of the presumptive actinomycete isolate JB66 was performed by molecular taxonomy using 16S rRNA gene sequence analysis. Genomic DNA was isolated as described previously. [38] The 16S ribosomal DNA of JB66 isolate was amplified employing universal primers 27F-5’AGAGTTTGATCTTGGCTCAG3’ and 1525-R 5’ AAGGAGGTATCCGCA 3’ using a Thermocycler. [44,45] The reaction was optimized with an initial denaturation at 95°C for 5 min, followed by 30 cycles of DNA denaturation at 95°C for 1 min, primer-annealing at 44°C for 1 min and extension cycle at 72°C for 1.5 min, with a final extension at 72°C for 10 min. PCR-amplicon was visualized in 2% agarose gel electrophoresis and subsequently revealed with ethidium bromide staining. The amplified 16SrDNA gene product was sequenced using the Sanger dideoxy method. Manual sequence edition, contig assembling and sequence alignment were executed using Vector NTI v10 software package. Sequencing results were analyzed for chimeras using DECIPHER v1.4.0 program. [46] The 16SrRNA gene sequences were compared with the GenBank/EMBL/DDBJ databases by using the BLASTN [47] search program. After pairwise alignment using CLUSTAL_X program v1.8 [48], the phylogenetic tree was constructed by using the neighbor-joining method [49] using MEGA X. [50] Kimura 2-parameter method was used to compute the evolutionary distances of JB66 with other members of the genera. [51]

**Detection of polyketide synthases-II gene**

A set of degenerate primers (KSα 5’-TSG CST GCT TGG AYG CSA TC-3’ and KSβ 5’-TGG AAN CCG CCG AAB CTT CT-3’) for the specific amplification of the gene encoding polyketide synthases-II (PKS-II) in the strain JB66 was used. [52-53] The PCR reaction mixture contained 2.5U of Taq DNA polymerase, 1mM MgCl2, 0.4mM deoxynucleoside triphosphates, 2 micromoles of each primer, and 5% dimethyl sulfoxide to a 50 microlitres volume. The actinobacterial DNA template was not added in the control reaction. The PCR reaction was optimized as one denaturation step of 94°C for 5 min, 30 amplification cycles of 94°C for 1 min, primer-annealing at 54°C for 1 min, and extension cycle at 72°C for 2 min and a final extension at 72°C for 5 min.

**Chemical screening of crude extract by thin-layer chromatography and bioautography**

The readymade pre-coated TLC plates (Merck; silica gel 60-F254 nm) were used for the separation of the methanolic crude extract of the strain JB66. Using a capillary tube, a row of spots of the active fraction of the extract was applied 1cm above the bottom of the TLC plate. The spots were then left to dry. The different solvent systems, including ethanol: water: chloroform (40:40:20), n-butanol: acetic acid: water (2:1:8), hexane: ethyl acetate (1:9), chloroform: methanol (18:2), dichloromethane: methanol (18:2), ethyl acetate: methanol (16:4), benzene: hexane (3:4) were initially used for the separation. Among different combinations, a mixture of chloroform: methanol (18:2) provided the best resolution and separation of chemical components from the JB66 methanolic extract. The TLC plate was placed vertically in a developing chamber containing the solvent chloroform: methanol in a ratio of 18:2, covered with the glass to prevent the evaporation. The solvent was allowed to migrate until it reaches 80% of the TLC plate. After that, the TLC plate was kept at room temperature for drying. After drying, the plate was viewed under UV (254 nm) and in iodine vapour.
The Rf value of the spot separated on the TLC plate was determined by the following equation:

\[
R_f = \frac{\text{Distance traveled by the solute}}{\text{Distance traveled by the solvent}}
\]

UV-visible absorption spectrum (200-600 nm) of the crude methanolic extract of JB66 was recorded on a spectrophotometer (Shimadzu) and bioautography was done using MDR pathogen *S. typhi*.

**RESULTS**

**Isolation of actinobacteria**

In the present study, many actinobacterial isolates were recovered from the samples collected from sand dune soil in the Bikaner district of the Thar desert in Rajasthan. Among various media tested, modified actinomycete selective (MAS) agar medium was found to be the best medium as it provided the highest yield of actinomycetes like colonies. An earthy smell, opaque, having a chalky appearance of fully grown colonies, strongly adhered to the agar medium, darker in the center, irregular, fuzzy edge, sometimes star-shaped, hyphal appearance and leathery look, putative actinomycete like colonies were selected.

**Screening for antimicrobial activity**

Based on primary and secondary screening, the isolate JB66 exhibited the broad-spectrum antimicrobial activity against the tested bacterial pathogens. The strain JB66 showed antimicrobial activity with the inhibition zones (measured in diameter) of 21 mm against *Shigella flexneri*, 20 mm against *E. coli*, 27 mm against *Staphylococcus aureus* and 20 mm against *Klebsiella pneumoniae*. Biological extracts (supernatant) collected from the JB66 strain exhibited the maximum growth inhibition of the pathogens tested in the present study after eight days of incubation and then declined (Figure 1).

**Biological activity of JB66 against MDR pathogens**

Two clinical isolates *P. vulgaris* and *S. aureus* were tested against various antibiotics (Hi-Media, Mumbai) to determine their antibiogram profile (Table 1).

Among the selected 20 antibiotics, *P. vulgaris* was found to be resistant (R) against 13 tested antibiotics namely amoxicillin-clavulanic acid (30µg), aztreonam (30µg), ceftriaxone (30µg), cefotaxime (30µg), cefuroxime (30µg), ertapenem (10µg), gatifloxacin (5µg), gentamicin (10µg), meropenem (10µg), netilmicin (30µg), norfloxacin (10µg), ofloxacin (5µg), piperacillin-tazobactam (100/10µg) antibiotics (Figure 2) and sensitive (S) against 7 antibiotics amikacin (30µg), cefepime (30µg), cefoperazone-sulbactam (75µg-30µg), ciprofloxacin (1µg), trimethoprim-sulfamethoxazole (25µg), imipenem (10µg), levofloxacin (5µg) with the zone of inhibition (in mm) 20, 20, 17.5, 20, 17, 23,19 respectively (Table 1). On the other hand, among 21 tested antibiotics, *S. aureus* was found to be resistant (R) against 16 antibiotics namely amoxicillin-clavulanic acid (30µg), ampicillin (10µg), azithromycin (30µg), cefepime (30µg), ceftriaxone (30µg), cefotaxime (30µg), cephalexin (30µg), cloxacillin (30µg), trimethoprim-sulfamethoxazole (25µg), erythromycin (15µg), imipenem (10µg), meropenem (10µg), piperacillin-tazobactam (100-10µg), teicoplanin (30µg), tetracycline (30µg), and vancomycin (30µg) antibiotics. While it was found sensitive against 5 antibiotics namely cefuroxime (30µg), ciprofloxacin (1µg), clindamycin (2µg), levofloxacin (5µg), linezolid (30µg) with zone of inhibition (in mm) 18, 21, 22, 20, 23 respectively (Table 1) (Figure 2). The methanolic extract of the isolate JB66 exhibited the growth inhibitory effect against both MDR pathogens by producing an inhibition zone diameter of 22 mm against MDR *S. aureus* and 19 mm against MDR *P. vulgaris* (Table 1) (Figure 3).

![Fig. 1: Analyses of antimicrobial product formation, optical growth density and growth inhibition by JB66 strain against E. coli, K. pneumoniae, S. flexneri and S. aureus.](image1)

![Fig. 2: Antibiotic susceptibility of MDR pathogens (i) S. and (ii) Proteus vulgaris. The antibiogram profile of pathogens was prepared using lcosa discs on a 200 mm plate. Both the pathogens were found resistant for most the antibiotics tested.](image2)

![Fig. 3: Antimicrobial activities of the methanolic crude extract from JB66 against multidrug-resistant (A) S. aureus (B) Proteus Vulgaris. JB66 strain exhibited the maximum growth inhibition against S. aureus measured as a diameter of inhibition zone (mm)](image3)
Morphological and physiological characteristics of JB66

The Gram-positive actinobacterial isolate JB66 showed typical mycelial structure in Gram's staining (Figure 4). The scanning electron micrograph of the isolate JB66 revealed a cubic form, smooth spore chain (Figure 4). It exhibited white-gray aerial mycelium and yellowish-white substrate mycelium on tryptone-yeast extract (ISP-1) agar, light gray aerial mycelium and pale white substrate mycelium on yeast extract-malt extract-dextrose (ISP-2) agar, light gray color aerial mycelium and pale white substrate mycelium on oat-meal agar (ISP-3), grey aerial mycelium and light brown substrate mycelium starch-inorganic salts (ISP-4) agar and grey aerial and yellowish white substrate mycelium on glycerol-asparagine (ISP-5) agar. The growth of JB66 was observed to be moderate on ISP6, exhibiting light grey aerial mycelium and grey white substrate mycelium whereas on tyrosine agar (ISP-7), it also showed moderate growth with light grey aerial and light brown substrate mycelium (Figure 5) (Table 2).

![Image](image-url)
moderate producer of cellulase, tyrosinase and urease enzymes while it was not observed to produce extracellular enzyme lipase. It showed positive activity for the catalase and gelatinase enzymes and the negative activity for DNAse enzyme.

Table 3: Biochemical and physiological characteristics

| Parameter       | Variables | Growth |
|-----------------|-----------|--------|
| Carbon Sources  |           |        |
| Arabinose       | +         |        |
| D-Mannitol      | +         |        |
| Galactose       | +         |        |
| D-Xylose        | +         |        |
| Sucrose         | +         |        |
| Rhamnose        | +         |        |
| Fructose        | +         |        |
| Dextrose        | +         |        |
| Lactose         | +         |        |
| Salicin         | +         |        |
| Meso-inositol   | +         |        |
| Ribose          | +         |        |
| Mannose         | +         |        |
| Pectin          | +         |        |
| D-Glucose       | +         |        |
| Alanine         | -         |        |
| Glycine         | +         |        |
| Histidine       | +         |        |
| Nitrogen Sources|           |        |
| Methionine      | -         |        |
| Glutamic acid   | -         |        |
| Aspartic acid   | +         |        |
| Proline         | +         |        |
| Tyrosine        | +         |        |
| Isoleucine      | +         |        |
| Cysteine        | +         |        |
| Tryptophane     | +         |        |
| Arginine        | +         |        |
| Asparagine      | +         |        |
| pH              |           |        |
| 5               | +         |        |
| 7               | +         |        |
| 9               | -         |        |
| 11              | -         |        |
| 25              | +         |        |
| Temperature (°C)|           |        |
| 37              | +         |        |
| 45              | +         |        |
| 55              | -         |        |
| NaCl (%)        |           |        |
| 0               | +         |        |
| 1               | +         |        |
| 2.5             | +         |        |
| 5               | +         |        |
| 7.5             | -         |        |
| 10              | -         |        |
| Amylase         | +         |        |
| Catalase        | +         |        |
| Cellulase       | +         |        |
| DNAse           | -         |        |
| Gelatinase      | +         |        |
| Extracellular enzyme |       |        |
| L-asparaginase  | +         |        |
| Lipase          | -         |        |
| Protease        | +         |        |
| Phosphatase     | -         |        |
| Tyrosinase      | +         |        |
| Urease          | +         |        |

+ good growth, - poor growth

Molecular characterization

The 16S rDNA sequence of the strain JB66 after Sanger sequencing was submitted to the GenBank database under an accession number MH762010. The query sequence was aligned and compared with the available 16S rDNA gene sequences in the GenBank database by using multi-sequence advanced BLAST comparison. It revealed that the isolate JB66 belongs to the genus *Streptomyces*. It showed 88.99-89.24% sequence similarity with its members and share the maximum (88.89%) similarity to *Streptomyces* sp. ATSC13. The phylogenetic analysis indicated that JB66 is clustered singly in the phylogenetic tree with the nearest neighbor exhibiting a similarity score of <97% (88.89%) suggesting the strain might belong to a possible new species (Figure 6).

Detection of PKS-II gene

To evaluate the biosynthetic potential, the strain JB66 was screened for the presence of PKS-II gene sequence by specific amplification of its chromosomal DNA.

![Fig. 6: The evolutionary relationship of JB66 was inferred using the Neighbor-Joining method. The % of replicate trees for the associated taxa (clustered for the bootstrap test) is shown next to the branches. The tree is drawn to scale. Kimura 2-parameter method was used to compute the evolutionary distances (number of base substitutions per site). Among sites, the rate variation was modeled with a gamma distribution (shape parameter = 2). Evolutionary analyses were conducted in MEGA X](image)

![Fig. 7: PCR amplification of PKS type II biosynthetic gene. An amplicon size of ~600bp, denoted by an arrow, for PKS-II gene, confirmed its presence in the actinomycete strain JB66](image)
Amplicon size of ~600bp for PKS type II confirmed the presence of this biosynthetic gene in the actinobacterial strain JB66 (Figure 7).

Fig. 8: Thin layer chromatograph showing the bio-active compound extracted from the fermentation broth of the JB66 strain. TLC was visualized under UV light and iodine spray.

Fig. 9: Bio-autography to assess the activity of various compounds produced by JB66 TLC. Rf values of 0.53 (possible cephalosporin like aromatic or polyene compound), 0.40 (possible demethoxy rapamycin like compound), 0.46 (possible daidzein like compound) fractionated compounds exhibited the strong inhibitory activity against S. typhi.

**Table 4: Putative compounds and their structures identified in the TLC profile of JB66**

| Rf | Name of compound                      | Structure of compound           |
|----|---------------------------------------|---------------------------------|
| 0.84 | Prodigiosin-like pigments            | ![Prodigiosin-like pigments](image) |
| 0.53 | Cephalosporin like aromatic or polyene compound | ![Cephalosporin like aromatic or polyene compound](image) |
| 0.46 | Daidzein like compound               | ![Daidzein like compound](image) |
| 0.40 | Demethoxy Rapamycin                 | ![Demethoxy Rapamycin](image)   |
| 0.33 | 4',6-Dihydroxy-7'-methoxyisoflavone | ![4',6-Dihydroxy-7'-methoxyisoflavone](image) |
| 0.03 | Amiclenomycin                        | ![Amiclenomycin](image)         |

**DISCUSSION**

The genus *Streptomyces* has gained global attention due to its ability to produce many useful biologically active metabolites. The most common source to isolate *Streptomyces* is soil; nevertheless, the rate of discovery of novel metabolites from broadly explored terrestrial strains has considerably diminished. [55] The probability of discovering novel metabolites could be increased by exploring the unexplored or less explored environments. Consequently, the bioprospecting of *Streptomyces* derived secondary metabolites from the less explored ecological niches has been recommended. [56-57] Therefore, an attempt was made in the present study to explore the unexplored Bikaner district of the Thar desert in Rajasthan to isolate actinomycetes producing therapeutically important metabolites. Although a few reports exist on the prevalence of actinomycetes in the desert ecosystems [11, 25, 58-63], to date, there has been no such a detailed study conducted to address the antimicrobial potential of these actinobacteria against multidrug-resistant pathogens. Among various isolates, recovered from the samples of sand dune soil, the isolate JB66 exhibited the broad-

**Thin-layer chromatography and bio-autography**

The separation of the metabolic compounds present in the methanolic extract of the strain JB66 was done using TLC. Among various solvent systems in various proportions, chloroform: methanol in 18:2 proportions showed a good separation. The Rf values of TLC fractionated compounds were calculated to be 0.83, 0.53, 0.46, 0.40, 0.33, 0.09, 0.03 (Figure 8). The Rf values of the separated spots on silica gel TLC led to the preliminary identification of prodigiosin pigment or chandramycin (Rf 0.84), cephalosporin or zeatin (Rf 0.53), daidzein (Rf 0.46), demethoxy rapamycin (Rf 0.40), 4,6-dihydroxy-7-methoxyisoflavone (Rf 0.33), munumbicins (Rf 0.09) and amiclenomycin (0.03) like compounds (Table 4). The bioautography of metabolites, fractionated on the TLC plate at different Rf values revealed that the fractions with Rf value of 0.40, 0.46, 0.53 value were actually associated with the antimicrobial activity. The inhibition zone around the excised TLC fraction (band/spot) showed the bioactive part against S. typhi (Figure 9). The JB66 extract showed UV-VIS spectrum absorbance maxima at 288 nm that indicated the presence of an aromatic nucleus.
spectrum antibacterial activity against the bacterial pathogens including MDR pathogens. Based on the phenotypic and molecular characteristics of the isolate JB66, it was identified as a member of the genus *Streptomyces*. [64] 16SrRNA gene sequence analysis revealed that it has the closest sequence similarity (~89%) with *Streptomyces* sp. (KT877000.1). The colour of the aerial and substrate mycelia of actinomycetes has been observed to be medium-dependant [65] and serves as one of the important colonial characteristics for taxonomical studies. [66–67] The isolates with abundant powdery spores and with an earthy odor of geosmin are usually classified as streptomyces-like strains. [68] Similarly, the actinomycete isolate JB66 grew well on the different ISP media used that further confirmed its identity as a member of the genus *Streptomyces*. The methanolic crude extract of the strain JB66 showed significant antimicrobial activity against MDR *P. vulgaris* and *S. aureus*. In a previous study, the in vitro potential of the bioactive crude extracts of five putative novel species of actinobacteria was assessed. These isolates were recovered from the Indian Thar desert that exhibited antibacterial activity against multidrug-resistant (MDR) *Streptococcus pneumonia*. [25] Saadoun and Gharaibehalso reported antibacterial activity of sixty different *Streptomyces* isolates from Badia Region of Jordan desert against *B. subtilis, S. aureus, E. coli, Klebsiella* sp. and *Shigella* sp. [69] The Selvameenal group also reported a yellow pigment producing *Streptomyces* from the Thar desert soil that exhibits antibiotic activity against methicillin-resistant *Staphylococcus aureus* (MRSA), vancomycin-resistant *Staphylococcus aureus* (VRSA) and extended-spectrum β-lactamases (ESBL) cultures of *E. coli* and *Klebsiella*, [62]

Thin-layer chromatography was used in the present study for the fractionation of crude methanolic extracts of the strain JB66. Among many of the solvent systems tested, chloroform: methanol (18:2) was found to be the most effective solvent system that provided the best separation. The different Rf values of the separated spots observed on the silica gel plate (TLC) led to the preliminary identification of various chemical compounds. The fraction with Rf value of 0.84 exhibited the presence of chondramycin like compound, previously this novel antibiotic was extracted with organic solvents from the culture broth of *Streptomyces halstedii-olivaceus* AA32 and AA33. [72]

Similarly, the fraction with Rf value of 0.40 exhibited the presence of demethoxy rapamycin like compound, earlier reported to be produced by *Streptomyces hygroscopicus* that posses a broad spectrum antagonistic activity against few Gram-positive and Gram-negative bacteria and fungi. [73] Rapamycin is a macrolycin natural product that exhibits various biological and pharmacological activities, including antifungal, immunosuppressive, antitumor, neuroprotective, and anti-aging activities. The fraction at Rf values of 0.46 and 0.33 exhibited the presence of isoflavonoids daidzein and 4,6-dihydroxy-7-methoxyisoflavone (Kakkatin) like compounds respectively. These are plant flavonoids that have been identified after chromatographic separation in *Streptomyces* sp. GW10/1811 and *Streptomyces* sp. GW39/1530. [74] The fraction with Rf value of 0.03 exhibited the presence of amiclenomycin like compound, which was observed to be produced by the culture filtrates of a *Streptomyces lavendulae* subsp. *amiclenomycini* (Streptomyces strain MD580-S1). The compound amiclenomycin has been reported to inhibit the growth of mycobacteria, including antibiotic-resistant mycobacteria. [75] The fraction with Rf value of 0.09 exhibited the presence of munumbcins like compound which was previously reported in *Streptomyces* NRRL 3052. The strain was isolated from medicinal plant snakevine (*Kennedia nigriscans*), native to the Northern Territory of Australia. This compound has been known to exhibit broad-spectrum activity against many human and plant pathogenic fungi, bacteria and *Plasmodium* sp. [76] In the present study, after the fractionation of crude extracts in TLC, the actual bioactive fractions were recognized using bio-autography. A total of three fractions of chemical compounds with Rf values of 0.40, 0.46, 0.53 of JB66 methanolic extract were observed to be bioactive against MDR pathogen. To confirm this further, putatively identified chemical compounds will be confirmed in LC-MS analysis. In conclusion, it can be stated that the metabolic compounds from the strain JB66 can serve as potential candidates for the development of chemotherapeutic drugs to control the infections caused by MDR pathogens in the future.

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