Functional Genome Screening to Elucidate the Colistin Resistance Mechanism

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Antibiogram profile of 1590 clinical bacterial isolates based on thirteen different antimicrobial compounds showed that 1.6% of the bacterial isolates are multidrug resistant. Distribution pattern based on 16S rRNA sequence analysis showed that *Pseudomonas aeruginosa* constituted the largest group (83.6%) followed by *Burkholderia pseudomallei* sp. A191 (5.17%), *Staphylococcus* sp. A261 (3.45%). Among the various antibiotics used, colistin appeared to be the most effective against the Gram negative bacteria. *Burkholderia pseudomallei* sp. A191 and *Pseudomonas aeruginosa* sp. A111 showed resistance to 1500 μg/ml and 750 μg/ml of colistin respectively which constitutes 7.7% of the bacterial population. A functional genomics strategy was employed to discover the molecular support for colistin resistance in *Burkholderia pseudomallei* sp. A191. A pUC plasmid-based genomic expression library was constructed with an estimated library size of 2.1 × 10⁷ bp. Five colistin resistant clones were obtained after functional screening of the library. Analysis of DNA sequence of five colistin resistant clones showed homology to two component regularity systems (TCRS) encoding for a histidine kinase (mrgS) and its regulatory component (mrgR). Cross complementation assay showed that mutations in mrgS were sufficient enough to confer colistin resistant phenotype in a sensitive strain.

Antimicrobial resistance (AMR) is a global public health crisis and is of great concern of every country irrespective of their socio-economic status1. Hospitals, more particularly intensive care units, are major sites of origin for the development and evolution of antibiotic resistant bacteria2. In developing countries, hospitals are crowded with debilitated patients who are administered with heavy doses of broad-spectrum antibiotics often without diagnosing specific infecting organism and without following any proper guidelines leading to an ineffective treatment. This brings in a threat to the patient survival and also to curb spread of infection3.

Outbreaks of infections with multi drug resistant (MDR) strains in intensive care unit settings have been reported in several countries across the world. The treatment of these infections has become difficult due to growing prevalence of pan drug resistance (PDR). Although, in these resistant strains, colistin (also known as polymyxin E) is often considered as the last resort of treatment, there are few reports on emergence of colistin resistance7. Few reports have been surfaced out in Indian subcontinent about the colistin resistance however the mechanism of its resistance has not been figured out so far8. Moreover, no report is available on the genetic mechanisms of colistin resistance in *Burkholderia* species.

In the present study, we investigated the mechanism of colistin resistance by functional genome screening in *Burkholderia pseudomallei*.

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Results
A total of 1590 bacterial isolates were collected from clinical samples (wound, burn injuries, sputum, pus, urine and head injuries) isolated from patients admitted to various wards of the hospital. Average age of the hospitalized patients was found to be 34 years of which 42.3% of the patients were female and 57.7% were male (Table 1).

Clinical isolates which showed resistance to more than any three antibiotics (1.6%) were selected for further study. Thirteen antimicrobial compounds belonging to various classes of antibiotics including β-lactam, quinolone, aminoglycoside, and polymyxin were used to generate the antibiogram profile. Within the β-lactam class, 92.3% of the isolates were resistant to amoxycllin and clavulanic acid followed by cefepime (84.61%) and ceftriaxone (53.8%) (Fig. 1). Among the aminoglycosides, 53.45% of the isolates showed resistance to gentamicin followed by netillin (34.6%). In the quinolone class, 46.15% and 34.61% isolates showed resistance to ciprofloxacin and oflloxacin respectively (Fig. 1). Colistin was found to be most effective because only 7.7% of the bacteria showed resistance to this antibiotic (Fig. 1).

| Isolates | Sex | Age | Source               |
|----------|-----|-----|----------------------|
| A51      | F   | 24  | wound                |
| A71      | M   | 22  | wound                |
| A81      | M   | 18  | wound-pus            |
| A101     | M   | 70  | wound                |
| A111     | M   | 52  | sputum               |
| A121     | M   | 60  | head injury          |
| A141     | M   | 52  | non-healing Ulcers   |
| A151     | F   | 75  | pus                  |
| A171     | M   | 44  | wound                |
| A181     | M   | 06  | urine                |
| A191     | F   | 20  | pus                  |
| A221     | M   | 22  | pus                  |
| A251     | F   | 20  | 60%burn              |
| A261     | F   | 20  | wound                |
| A271     | F   | 20  | wound                |
| A281     | M   | 58  | pus                  |
| A291     | M   | 26  | urine                |
| A301     | F   | 50  | pus                  |
| A311     | M   | 40  | pus                  |
| A321     | F   | 27  | urine                |
| A331     | F   | 14  | pus                  |
| A341     | M   | 40  | pus                  |
| A351     | M   | 50  | pus                  |
| A361     | F   | 66  | urine                |
| A371     | F   | 57  | fish eye             |
| A381     | M   | 70  | traumatic burning injury |

Table 1. Information of the patients and source of sampling of clinical bacterial isolates.

Figure 1.  Antibiogram of the selected isolates.
Identification of colistin resistant clinical isolates through 16S rRNA sequence analysis revealed predominance of *Pseudomonas aeruginosa* constituted (83.6%). Sequence similarity analysis clustered *Pseudomonas aeruginosa* into seven subgroups which were named as (GpI–GpVII) (Fig. 2). GpI showed 99% similarity to *Pseudomonas aeruginosa* NO2, GpII was 97% similar to *Pseudomonas aeruginosa* strain PA5-1-2, GpIII showed 98% similarity to *Pseudomonas aeruginosa* strain HK1-2, GpIV was 98% similar to *Pseudomonas aeruginosa* strain T1, GpV showed similarity to *Pseudomonas aeruginosa* strain VRKPC5, GpVI was 97% similar to *Pseudomonas aeruginosa* strain D2 and GpVII showed 98% similar to *Pseudomonas aeruginosa* strain NBAII AFP-7. Other groups identified were *Burkholderia pseudomallei* sp. A191 (5.17%), *Staphylococcus* sp. A261 (3.45%), *Micrococcus* sp. A171 (2.58%), *Aeromonas* sp. A341 (2.58%) and *Acinetobacter* sp. A341 (2.58%).

It was alarming to find that *Burkholderia pseudomallei* sp. A191 and *Pseudomonas aeruginosa* sp. A111 (GpII) showed resistance to higher concentration of colistin ie. 1500 μg/ml and 750 μg/ml respectively. MIC value of colistin for *B. pseudomallei* 1026b was reported as 128 mg/ml. Cell free extracts of both the isolates however, did not show any enzymatic degradation of colistin (Table S1). Functional genomics library of *Burkholderia pseudomallei* sp. A191 and *Pseudomonas aeruginosa* sp. A111 were of size of 2.1 × 10⁷ bp and 1.30 × 10⁶ bp respectively.

Five colistin-resistant clones were obtained after functional screening of *Burkholderia pseudomallei* sp. A191, but no resistant clone was observed in *Pseudomonas aeruginosa* sp. A111 library. Interestingly, after analysis of insert DNA sequences from all the five *Burkholderia pseudomallei* clones, it was observed that all clones have a common DNA sequence or gene that could be responsible for conferring colistin resistance. Based on ORF prediction, two components regulatory system encoding for a histidine kinase (mrgS) and its regulatory component (mrgR) were observed in the clone DNA sequence that showed 98–99% similarity with other histidine kinase sequences of *Burkholderia pseudomallei*. Histidine kianse (mrgS) was PCR amplified, sub-cloned and sequenced (Fig. 3A). Six point mutations were observed in mrgS gene viz. V143M, P246R, G695A, G696R, R1048H and R1072C (Fig. 3C). Resistance model based on two component regulatory system which controls the expression of genes responsible for LPS modification has been proposed (Fig. 3D). Colistin sensitive strain of *Burkholderia pseudomallei* was not available with us thus other Gram negative colistin sensitive species were explored for complementation assay. Transformation of *Pseudomonas aeruginosa* sp. A71 with mrgS plasmid causes the development of colistin resistant phenotype. Microdilution assay showed that *Pseudomonas aeruginosa* sp. A71+ mrgS plasmid was able to grow with varying concentrations of colistin (≥100 μg/ml) as compared with *Pseudomonas aeruginosa* sp. A71 transformed with vector alone (Fig. 3B and Table 2).

**Discussion**

The burden of infectious diseases in India is the highest in the world which gets further aggravated due to the inappropriate and irrational use of antimicrobial agents against these diseases, resulting in increased incidence of development of antimicrobial resistance. Surveillance of antimicrobial resistance in microbial populations associated with patients is not conducted in hospitals due to lack of proper infrastructure. Lack of information about the antimicrobial resistance patterns in patients presents an obstacle to disease management, with respect to antimicrobial therapy, patient prognosis, and infection control. 16S rRNA sequencing of sample isolates to identify the diverse multidrug resistant bacteria such as *Burkholderia pseudomallei* sp. A191, *Micrococcus* sp. A71,
Aeromonas sp. A341, Staphylococcus sp. A261 and different strains of Pseudomonas aeruginosa acted as an alternative to standard classical tools and to prevent the misidentification of clinically relevant isolates. All the bacteria identified in this study were found to be resistant to multiple classes of antibiotics though colistin was tested as the most effective drug against these bacteria. Among the isolates, Burkholderia pseudomallei sp. A191 and Pseudomonas aeruginosa sp. A111 showed high level colistin resistance (>500 μg/ml). Burkholderia species is known to be intrinsically resistant to colistin but whichever few reports of colistin resistance have been emerged out are centered on Pseudomonas aeruginosa and Klebsiella pneumoniae. It is an alarming situation and clinicians should be aware that colistin resistance can occur in P. aeruginosa, and some of these strains have the capability for cross contamination within a hospital set up. Occurrence of multi drug resistance in P. aeruginosa and capability of P. aeruginosa for cross contamination within a hospital set up is quite alarming to clinicians and also research scientist working in this field. Lee and Ko tried to correlate the two component regulatory systems PmrAB and PhoPQ with colistin resistance in Pseudomonas aeruginosa. However no amplification was obtained using the primers specific for TCRS based on Pseudomonas aeruginosa PA14 which could be attributed to difference with in the genomes at strain level (Data not shown). Alternatively, may be a different mechanism is prevailing in this bacterium responsible of resistance which is yet to be clearly identified. However, no reports on the mechanism of colistin resistance in Burkholderia sp. are available. Various major and minor determinants of polymyxin B like; truncation of the LPS core oligosaccharide, sigma factor RpoE, zinc metalloproteases and an efflux system (NorM) have been proposed to be responsible for bringing resistance in Burkholderia sp. Among these determinants, two component response regulator (BCA2831) and a periplasmic protease (MucD) are less studied in Burkholderia cenocepacia. Lack of potential of cell free extracts in catalytic degradation of colistin ruled out the role of protease in colistin conversion. A
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Table 2. Cross complementation of mrgS mutations.

| Strain                          | Colistin resistance (μg/ml) |
|---------------------------------|-----------------------------|
| Pseudomonas aeruginosa A71 + vector | <0.5                        |
| Burkholderia pseudomallei A91  | ≥100                        |
| Pseudomonas aeruginosa A71 + mrgRS | ≥100                        |
stimuli like pH or metal ions trigger the activation of TCRS\(^1\). Functional genome screening led us to discover the TCRS from *Burkholderia pseudomallei* sp. A191 and sequence analysis revealed unique mutations in *mrgS*. Since DNA sequences in all the clones were same, we hypothesized that mutations within the TCRS may result in the constitutive activation and subsequent expression of LPS modifying genes (Fig. 3D). This modification is carried out by the products of the *pmrHFIJKLM* operon, which is conserved among Enterobacteriaceae and is positively regulated by the PhoQ/PhoP and PmrAB signaling systems\(^2\).

To confirm this hypothesis, cross complementation assay were initiated which showed that mutations in two component regulatory system *mrgRS* of *Burkholderia pseudomallei* sp. A191 were sufficient to confer colistin resistance in the sensitive strain of *Pseudomonas aeruginosa* sp. A71. Non availability of colistin sensitive *Burkholderia* sp. forced us to do complementation assay with *Pseudomonas aeruginosa* sp. A71. To date it has not been possible to isolate the *Burkholderia* sp. which showed sensitivity to polymyxins\(^6\). Cross complementation assay demonstrated the possibility of dissemination of colistin resistant determinant among the clinical isolates within a hospital.

It can be concluded that regular surveillance to track the isolates with correct identification is highly important to tackle the antibiotic resistant pathogens. Discovery of colistin resistance at such a high concentration is alarming which highlights the need for strict measures to keep a tab on judicious antibiotic usage for infection control. Genetics of colistin resistance showed that single gene mutations are sufficient to confer the resistance. Cross complementation assays demonstrated that spread of resistant determinants is highly possible within a hospital.

**Methodology**

**Sample collections.** Institute of Medical Sciences and Sum Hospital (IMS & SUM), Siksha “O” Anusandhan University, Bhubaneswar, Odisha, is a more than 1000 bed hospital and handles 1500 patients on daily basis. Samples were collected from hospitalized-patients admitted to different wards of hospital during August 2012 to September 2013 (Table 1). Primary identification of bacteria was done based on the standard conventional morphological and biochemical tests.

**Ethical Considerations.** All participants were recruited under informed consent form in accordance with the approved guidelines from “Indian Council of Medical Research”, signed either by the patient or their family member. All experimental protocols were approved by the Institutional Ethical Committee, Center of Biotechnology (School of Pharmaceutical Sciences), SOA University, India. All the experimental methods were carried out in accordance with the approved guidelines.

**Antimicrobial susceptibility testing.** The susceptibility of these isolates was tested against antimicrobial agents according to the Clinical and Laboratory Standards Institute (CLSI) guidelines (2015). End-points were read after overnight incubation at 37 °C. The test microbes were taken from broth culture with inoculating loop and transferred to test tubes containing 5.0 ml sterile distilled water. The inoculums were added until the turbidity became equal to 0.5 McFarland standards. Cotton swab was then used to inoculate the test tube suspension onto the surface of the Muller Hinton agar plate and the uniformly swabbed plates were then allowed to dry. Antimicrobial agents and ranges (\(\mu g/ml\)) tested were: amoxycillin/clavulanic acid (4–32), ciprofloxacin(0.125–8), ceftriaxone (8–64), amikacin (1–64), ceftazidime (16–32), cefepime (2–32), gentamicin (2–16), netin (4–32), ceftoperazone (0.05–64), tobramycin (2–16), ofloxacin (2–8), imipenem (0.5–16), Cefpirome (0.25–4) and colistin (0.125–32). *P. aeruginosa* ATCC 27853 was used as a control strain.

**Molecular identification of antibiotic resistant bacteria.** Chromosomal DNA of the bacteria was extracted as described by Kumar et al.\(^{22}\). 16S rRNA gene was PCR amplified using genomic DNA as template with universal primers E8F (AGAGTTTGTATCCTGCGCTCACG) and reverse primer E1492R (GGT-TACCTTGTTACGACTT)\(^{24}\). The PCR reactions were performed as described by Kumar et al.\(^{22}\) and thermal cycling was performed as described by Kumar et al.\(^{22}\). Sequencing was done by Amnion Biosciences, Bangalore, India and then BLAST searched through the NCBI GenBank database. Phylogenetic tree was constructed using molecular evolutionary genetics analysis (MEGA) software with 1000 bootstrap replicates\(^25\).

**Enzymatic assay.** Cell extracts were incubated with colistin (100\(\mu g/ml\)) in a buffer. After regular interval of time, samples were analyzed by HPLC for the estimation of degradation of colistin. HPLC Prominence system (Shimadzu, Singapore) equipped with a binary LC-20AD pumping system with an online vacuum degasser, SIL-20 autosampler, and SPD-M20A photodiode array detector (PDA) detector was applied to chromatographic studies. Chromatographic separations of colistin were achieved on the C18 column (150 mm \times 4.6 mm i.d.), Phenomenex, USA. Samples were filtered through a Millipore membrane (0.45 \(\mu m\)). LCsolution software was used for data acquisition and integration. The UV detector was set at 214 nm and the temperature was ambient temperature. The sample injection volume of the autosampler was 5.0 \(\mu l\). The chromatographic separation was performed using a linear gradient (20% B (acetonitrile)+80% A (0.05% TFA aqueous solution) changed to 50% B+50% A in 10 min).

**Genomic expression library construction.** A genomic expression library was constructed by shearing the genomic DNA (gDNA) in order to obtain 7.0–9.0 kb DNA fragments after agarose gel electrophoresis. The sheared DNA fragments were end- repaired and ligated into a pUC-19 plasmid and were then electrotransformed into colistin sensitive *Pseudomonas aeruginosa* sp. A71 as host. Insert size distribution was estimated by gel electrophoresis of colony PCR products obtained by amplifying the insert using vector specific primers. The total size of the genomic expression library was determined by multiplying average PCR based insert size by the number of total CFUs obtained. The transformation mixture was enriched by growing the cells in selective LB broth.
and glycerol preserved at −70°C until further processing. Resistant clones containing unique DNA inserts were sequenced using Sanger sequencing technology (Amnion Biosciences, Bangalore, India).

**Molecular cloning of two component regulatory systems.** Recombinant plasmid was extracted from a resistant clone. Oligonucleotides were designed to amplify the two component regulatory system (TCRS) mrgRS coding for transcriptional regulator and kinase sensor using plasmid as template.

B.mrg-R 5'ATGTCCACGTTGCCCTGAATC3'
B.mrg-F 5'CATAGCGATCTTATGCGGCA3'
B.mrgS-F 5'ATGCAAAGCCTCCTGAAGA3'
B.mrgS-R 5'TCACGGCGGGCGGCTTTCC3'

PCR amplification was performed as described above. Amplicon was ligated into pTZ57R vector (pUC based) as per manufacturer's instructions and transformed into electrocompetent *Pseudomonas aeruginosa* A71.\(^2\) The presence of this plasmid in the colonies was confirmed using PCR with universal M13 primers where the amplification of the product in this PCR required the junction between the mrgRS sequence and the vector to be present. The sequence homology search and conserved domain analysis of the deduced protein sequence were performed using the BLASTP program (http://blast.ncbi.nlm.nih.gov/Blast.cgi) and CDART program (http://www.ncbi.nlm.nih.gov/Structure/lexington/lexington.cgi) of NCBI, respectively. The ORFs were identified by using the NCBI's open reading frame (ORF) Finder tool (http://www.ncbi.nlm.nih.gov/entrez/query.fcgi).

**Cross complementation assay.** To determine whether the mutations in mrgS were sufficient to confer colistin resistance, a broth microdilution assay was used to determine the MIC of sensitive strain *Pseudomonas aeruginosa* A71 carrying the empty vector or the vector with the mrgS gene. Overnight culture of *Pseudomonas aeruginosa* A71 grown in Mueller-Hinton broth (MHB) plus carbencillin (200 μg/ml) was harvested by centrifugation, washed with phosphate buffer, re-suspended in MHB broth, and inoculated into MHB with a series of colistin concentrations.

**Nucleotide sequence submission.** The nucleotide sequences of 16S rDNA of the clinical bacterial strains reported in this study was deposited in the GenBank database with the following accession numbers *Acinetobacter baumannii* (KT819271), *Micrococcus sp.* A171 (KT819272), *Burkholderia pseudomallei* A191 (KT819273), *Staphylococcus* sp. A261 (KT819274), *Aeromonas* sp. A341 (KT819275), *Pseudomonas aeruginosa* A361 (KT819276), *Pseudomonas aeruginosa* A111 (KT819277), *Pseudomonas aeruginosa* A151 (KT819278), *Pseudomonas aeruginosa* A301 (KT819279), *Pseudomonas aeruginosa* A311 (KT819280), *Pseudomonas aeruginosa* A71 (KT819281), *Pseudomonas aeruginosa* A81 (KT819282).

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Author Contributions
M.K. designed all the research, performed the experiments, analyzed the data and wrote the manuscript. A.G., R.K.S. and J.J. performed the experiments. N.K.D. and E.S. helped to write the manuscript. All authors reviewed and approved the manuscript.

Additional Information
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