Identification of Novel Regulatory Small RNAs in Acinetobacter baumannii

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Abstract

Small RNA (sRNA) molecules are non-coding RNAs that have been implicated in regulation of various cellular processes in living systems, allowing them to adapt to changing environmental conditions. Till date, sRNAs have not been reported in Acinetobacter baumannii (A. baumannii), which has emerged as a significant multiple drug resistant nosocomial pathogen. In the present study, a combination of bioinformatic and experimental approach was used for identification of novel sRNAs. A total of 31 putative sRNAs were predicted by a combination of two algorithms, sRNAPredict and QRNA. Initially 10 sRNAs were chosen on the basis of lower E-value and three sRNAs (designated as AbsR11, 25 and 28) showed positive signal on Northern blot. These sRNAs are novel in nature as they do not have homologous sequences in other bacterial species. Expression of the three sRNAs was examined in various phases of bacterial growth. Further, the effect of various stress conditions on sRNA gene expression was determined. A detailed investigation revealed differential expression profile of AbsR25 in presence of varying amounts of ethidium bromide (EtBr), suggesting that its expression is influenced by environmental or internal signals such as stress response. A decrease in expression of AbsR25 and concomitant increase in the expression of bioinformatically predicted targets in presence of high EtBr was reverberated by the decrease in target gene expression when AbsR25 was overexpressed. This hints at the negative regulation of target genes by AbsR25. Interestingly, the putative targets include transporter genes and the degree of variation in expression of one of them (A15_1331) suggests that AbsR25 is involved in regulation of a transporter. This study provides a perspective for future studies of sRNAs and their possible involvement in regulation of antibiotic resistance in bacteria specifically in cryptic A. baumannii.

Introduction

During the last three decades, nosocomial infections caused by multidrug resistant (MDR), Gram-negative bacilli have become a significant health problem. A. baumannii is a versatile Gram-negative bacillus that has evolved to be one of the major opportunistic nosocomial pathogens [1,2]. It causes bloodstream infections, ventilator-associated pneumonia, urinary tract infections, meningitis and wound infections in hospitalized patients; especially the ones with prolonged stays in intensive care units where the host is most vulnerable and the use of antimicrobial agents is highest [3,4]. A. baumannii is a strictly aerobic non-motile bacterium that can adhere to the inanimate surfaces like the hospital equipments and survive on these dry surfaces for long periods of time [5–7]. Difficulties in controlling and eradicating the spread of A. baumannii have challenged clinicians worldwide. Its intrinsic resistance to a range of antimicrobial compounds, ability to acquire antibiotic resistance genes by horizontal transfer and global spread has led to high mortality in patients infected with this organism [1,8–10].

Resistance of A. baumannii to almost every clinically used antibiotic makes treatment of infections extremely difficult [11]. Multidrug resistance in A. baumannii is attributed to different mechanisms including degradation, modification, decreased permeability and increased efflux of antibiotic molecules [12]. Multidrug efflux pumps that decrease drug accumulation in the cell, cause severe complications in the treatment of infections caused by A. baumannii. Several efflux systems conferring resistance to antimicrobial agents have been described in A. baumannii [13–18] and many putative efflux pump genes have been identified in its genome sequence [19]. To understand how the bacterium utilizes these efflux pumps, it is important to first explicate the process by which their expression is regulated. The study of the intrinsic regulation of expression of these pumps presents a viable option to elucidate the behaviour and virulence of the pathogen and thus design therapies directed against the pathogen.

Regulation by RNA (Ribonucleic Acid) molecules plays a significant role in amending gene expression in all living systems including bacteria. The largest and most extensively studied bacterial regulatory RNAs are non-coding RNAs commonly
referred as small RNA (sRNA) due to their small size (50–500 nucleotides) [20,21]. sRNAs control the expression of genes by base pairing to their cognate target mRNA and modulating the translational activity and/or the stability of mRNA. sRNAs are significant to a pathogen as they regulate virulence [21–26] and susceptibility to different stress conditions [27–30]. Initially, sRNAs were identified in *Escherichia coli*, however, with increase in availability of bacterial genome sequence, many sRNAs have been identified in pathogenic bacteria [31] like *Mycobacterium tuberculosis* [32], *Stenotrophomonas* spp. [33–35], *Listeria monocytogenes* [36], *Pseudomonas* spp. [36–39], *Vibrio* spp. [40,41] and *Staphylococcus* spp. [42–45]. A number of these sRNAs have been implicated in the regulation of genes significant for bacterial drug resistance [46–49].

Although it is widely accepted that sRNAs are encoded by most if not all bacteria, genome-wide annotations for sRNA-encoding genes have not been conducted in the clinically significant pathogen, *A. baumannii*. Due to the importance of sRNAs as regulators, it is likely that they may have vital regulatory roles in drug resistance of *A. baumannii*. In the study reported here, a systematic search for sRNAs focussing on intergenic regions has been undertaken in *A. baumannii*. This search involved computational approaches followed by verification by Northern blotting leading to the novel sRNAs that we refer to as AbsR1 to 31 (for conditions). This study demonstrates for the first time the presence of sRNAs in the nosocomial pathogen *A. baumannii* and the expression of AbsR25 in different stress conditions.

**Materials and Methods**

**Sequence data and databases**

ORF databases, tRNA and rRNA coordinates were obtained from the national center for biotechnology information (NCBI) ftp database (ftp://ftp.ncbi.nih.gov/genomes/Bacteria/). Locus and product names were already mentioned in the database obtained from NCBI. Genomes of all the bacteria used in the present study were downloaded from NCBI, in FASTA format for basic local alignment search tool (BLAST) analysis.

**Prediction of sRNA loci using bioinformatic tools**

Candidate sRNAs in *Acinetobacter baumannii* ATCC17978 were predicted using the sRNApredict program [50] and other bioinformatic tools listed below:

**AB-BLAST.** This is a new version of BLAST developed by the Advanced Biocomputing Company based on the previous WU-BLAST developed by Warren R. Gish for Washington University. For AB-BLAST against *Pseudomonas aeruginosa*, *Burkholderia multivorans* and *Stenotrophomonas maltophilia*, the parameters were set as follows: word length (W) 7, match/mismatch scores (M/N) 5/4, penalty score for extending a gap (R) 10 and penalty for a gap of length one (Q) 10, and maximum E-value 1e-05.

These parameters correspond to default parameters of WU-BLAST 2.0, the software used in the study of *P. aeruginosa* sRNAs by Jonathan Livny et al [50]. The same parameters were used for BLAST against the genomes of *Acinetobacter* spp. (*Acinetobacter baylyi* ADP1) and *Acinetobacter* DR1 in a parallel study.

**QRNA.** It is a probabilistic model for predicting conserved secondary structure and its best probable fit. This program has three models, one for each of RNA, coding region and something else (for others). So, given a pairwise sequence alignment, it calculates Bayesian probability whether the sequence should correspond to an RNA, a coding region or something else. This is given in the output as ‘winner = <RNA/COD/OTH>’ for RNA, coding region or other respectively [51,52]. For QRNA script ‘blastn2qrnadepth.pl’ the parameter for max e-value was set to 1e-05. Further, for the QRNA program, the parameter for window span was set to 150 for database of *P. aeruginosa*, *B. multivorans* and *S. maltophilia* and 100 for database of two *Acinetobacter* spp. genomes and the slide size was 50 in both cases. Also, in the case of two *Acinetobacter* spp., the minimum length was set to 50 base pairs.

**RNAmotif.** This program takes a fasta formatted sequence file and a descriptor file, which gives the program a description of the key features of the structure to be looked for in the sequences, as input. For our purpose, the descriptor file gave the description of the key features of a rho-independent terminator [53]. This descriptor file was provided by D.J. Ecker, the creator of the program, to Jonathan Livny for his study of sRNAs of *P. aeruginosa* [50] and *Vibrio cholerae*. This file was available with the sRNApredict3 program as it can take input directly from RNAmotif if this file is used as descriptor file.

**TransTermHP.** It is a web-based software for predicting rho-independent terminators in a genome [54]. For our purpose, we kept the default settings except for the confidence level of the predicted terminators which was set to 96%.

**IGRExtract.** This program takes the fasta format file of the whole genome as an input along with the Genbank file and .ptt file for protein coding regions [55]. From these, it calculates the coordinates of inter genomic regions and extracts the sequences from the fasta file.

**sRNApredict3.** It is a C++ program that predicts sRNA using a coordinate based algorithm [55]. This algorithm requires the results of all the programs listed above to identify sRNA by searching for co-localization of genetic features predicted by these programs. These features have been generally known to be associated with genes encoding for sRNA such as rho-independent terminators (from RNAmotif and TransTermHP), secondary structure conservation (from QRNA), etc. The rRNA, tRNA and ORF loci were excluded by the sRNApredict from its output, which were downloaded from NCBI database.

**Bacterial strains and growth conditions**

*A. baumannii* MTCC1425 (ATCC15308) and *E. coli* DH5α were the bacterial strains used for this study. The strains were grown on nutrient agar (HiMedia, India) and Luria Bertani (LB) agar (Merck, Germany) medium respectively. For liquid cultures, *A. baumannii* was grown in nutrient broth at 30°C, 200 rpm and *E. coli* was grown in LB broth at 37°C under agitation (200 rpm). pTZ57R/T cloning vector (Fermentas, USA) was used for cloning RACE PCR products. Whenever needed, ampicillin was added to the LB broth at a final concentration of 100 μg/ml.

**Plasmid and strain construction**

Standard methods were used for plasmid isolation, DNA purification, restriction endonuclease cleavage, ligation and transformation [56]. Over-expression of AbsR25 was achieved under the control of arabinose inducible promoter P*BAD*, pPROBE-NT [57], a broad host range promoter less plasmid, was used for cloning the promoter (P*BAD*) and AbsR25. The plasmid was modified for replication in *A. baumannii* by cloning 1337 bp ori region from pWH1266 [58], a *Acinetobacter* genus specific plasmid, in EcoRI restriction site of pPROBE-NT. P*BAD* promoter containing araC, required for tight regulation, was
amplified form pBAD30 plasmid and AbsR25 was amplified from purified chromosomal DNA of A. baumannii MTCC: 1425. PBad promoter was cloned within SacI and XhoI enzyme sites of the plasmid and AbsR25 was cloned downstream of the promoter within SacI and HindIII sites. After each ligation the ligated product was transformed into E. coli DH5α cells and grown on LB plates containing 40 μg/ml kanamycin. Plasmids were recovered from the positive transformants and confirmed by digestion with respective enzymes. These cloning steps resulted in plasmid pPROBE-PBad-absR25 expressing absR25 under the control of arabinose inducible promoter. Primer sequences used are listed in Table 1. Furthermore the plasmid was sequenced and transformed into A. baumannii MTCC 1425 cells and selected by growth in LB agar plates containing 40 μg/ml kanamycin. 2% L-arabinose was used for induction of the promoter for 5 hrs.

Total RNA isolation and Northern blot analysis

For total RNA isolation, a single colony of A. baumannii MTCC1425 was grown overnight in a nutrient broth medium at 30°C. The culture was diluted to 1:100 times and grown at 30°C in 250 ml flasks with shaking at 200 rpm. Cells were harvested from culture grown to an optical density at 600 nm (OD600) of 0.4 (lag phase), 1 (early exponential phase), 1.6 (exponential phase) and 3.2 (stationary phase). Temperature, sodium chloride (NaCl) and ethidium bromide (EtBr) stresses were applied to the cells when the A. baumannii culture reached an OD600 nm of 0.4. For the temperature stress, the culture was incubated at different temperatures for 30 min and for NaCl stress, cultures were incubated with different concentrations of NaCl for 30 min. EtBr stress was provided by incubating culture with different concentrations of EtBr. Total RNA was isolated with TRI reagent (MRC, Cincinnati, Ohio, USA) for two hrs at 42°C and was pre-hybridized in ULTRAhyb buffer (Ambion, Austin, TX, USA) for two hrs at room temperature for two times, 5 minutes each and then with high stringency wash buffer (Ambion, Austin, TX, USA) for two times, 10 minutes each at 42°C according to manufacturer’s instructions. Following washes, the blots were developed (Bright Star Bio-detection kit, Ambion, Austin, TX, USA) and exposed to X-ray film for their visualization.

$5'$ and $3'$ rapid amplification of cDNA ends ($5'$ and $3'$ RACE)

$5'$ RACE was carried out on DNA free total RNA isolated previously, using the First Choice RLM-RACE kit (Ambion, Austin, TX, USA) according to the manufacturer’s instructions with slight modifications. Call Intestinal Phosphatase (CIP) treatment at the beginning was omitted, because prokaryotic RNA was used and a control without Tobacco Acid Pyrophosphatase (TAP) treatment was included. The AbsR specific primers (Table 1) were used in PCR steps of $5'$ RACE protocol and Hot star DNA polymerase (Qiagen, USA) was employed for amplification of the sRNA genes. The $5'$ RACE PCR product was gel purified and cloned into pTZ57R/T cloning vector (Fermentas, USA). The potential clones were selected, the plasmid DNA was isolated and sequenced for identification of transcription start site.

For $3'$ RACE, the total RNA was polyadenylated using Poly (A) Polymerase enzyme (New England Biolabs, USA) according to the manufacturer’s instructions. Polyadenylated RNA was then reversed transcribed and PCR amplified by using First choice RLM RACE kit (Ambion, Austin, TX, USA). The PCR product was cloned into pTZ57R/T cloning vector (Fermentas, USA) and sequenced for identification of the transcription termination site.

Target prediction of AbsR25 sRNA

In order to predict target mRNA genes for the identified AbsR25, sRNATarget [http://ccb.bmi.ac.cn/srnatarget/prediction.php] [59] and IntaRNA [http://rna.informatics.unifreiburg.de:8080/IntaRNA.jsp] [60] servers were used. Target genes with the score 1 in sRNAtarget were used for comparison analysis with IntaRNA output. For sRNAtarget, the default parameters were used. The default parameters for IntaRNA were as follows: Exact number of base pairs in seed: 7, folding window size: mRNA: 140, maximum base distance: mRNA: 70, maximum energy of interaction: 0 kcal/ml.

Quantitative Real Time-Polymerase Chain Reaction (qRT-PCR)

qRT-PCR was performed for detection of expression of bioinformatically predicted putative targets of AbsR25 sRNA. For sample preparation, a single colony of A. baumannii was grown overnight in nutrient broth at 30°C. This culture was diluted 100 times in 10 ml nutrient broth medium and grown till OD600 reached 0.4. When OD600 reached the desired point, the cultures were stressed with different concentrations 0, 64, 128, 192 and 256 μg/ml of EtBr and grown for 18 hrs at 30°C with shaking. After incubation, cells were pelleted and total RNA was isolated from each sample with TRI reagent (MRC, Cincinnati, Ohio, USA) according to manufacturer’s instructions, followed by DNase I (Fermentas, USA) treatment for removal of contaminating genomic DNA. Reverse transcription was performed for each stressed sample with Moloney Murine leukemia Virus reverse transcriptase (Promega, Madison, USA) and random hexamer primer. Negative control reactions were performed using equal concentrations of RNA but without Reverse transcriptase. Primers for Real Time PCR were designed with Primer3 software and are listed in Table 1. 16S rRNA was used as internal control for normalization of target mRNA gene expression. All cDNA aliquots for each stress condition were used in equal concentration in a qRT-PCR reaction mix containing 12.5 μl LightCycler 480 SYBR Green I master mix (Roche diagnostics, Germany), 0.2 μM each primer (forward and reverse) and reaction volume was made up to 25 μl with nuclease free water. The PCR was run on a
Smart Cycler (Cepheid, Sunnyvale, CA) with an initial denaturation for 5 min at 95°C and a subsequent run of 35 cycles, each comprising 15 sec at 95°C, 30 sec at 50°C and 30 sec at 72°C. The Ct (threshold cycle) values from all RT-PCR reactions in triplicate were analysed to detect target mRNA genes expression.

AbsR25 was over-expressed under the control of arabinose inducible promoter and qRT-PCR was used to analyze the effect of over-expression of AbsR25 on target genes. For sample preparation, a single colony of A. baumannii harbouring recombinant plasmid, pPROBE-PBAD-absR25, was grown overnight in nutrient broth at 30°C. This overnight culture was diluted 100 times in 10 ml nutrient broth medium and grown till OD 600 reached 0.6. When OD 600 reached the desired point, the culture was induced with 2% L-arabinose and grown for 5 hrs at 30°C with shaking. After incubation, cells were pelleted and total RNA was isolated from each sample with TRI reagent (MRC, Table 1. Oligonucleotides used in this study.

| Oligonucleotide | Sequence | Description |
|-----------------|----------|-------------|
| AbsR11_5’_RACE | ACA ACC TCG GTG AAG AGT CCC | For RACE mapping |
| AbsR11_3’_RACE | GAGAAGTCTAATTTTAAACTAGCG | For RACE mapping |
| AbsR25_5’_RACE | GCTTTTTTAACTTTAGCATCG | For RACE mapping |
| AbsR25_3’_RACE | TTTAAGCTTCTTAAATATCATGTG | For RACE mapping |
| AbsR28_5’_RACE | CGGAAGATGAAATTATTTGTAC | For RACE mapping |
| AbsR28_3’_RACE | CAAGTAAACACAGTAAACG | For RACE mapping |
| ABFC3251       | GCCAATAGCTCCTCCCAAAAGA | primer for qRT-PCR |
| ABFC3251       | TTCTCGACTTGGTGAAACCG | primer for qRT-PCR |
| ABFP1791       | TAGAATGTCGCGAAGACAGC | primer for qRT-PCR |
| ABFP1791       | TGCAAGAAGTCGACAAAGT | primer for qRT-PCR |
| ABFP2269       | GCAATATTGTGAATGCGCCC | primer for qRT-PCR |
| ABFP2269       | GTGGTATGAGTGAATGCGA | primer for qRT-PCR |
| ABFP2584       | CGCAACCAAGAAGATGAA | primer for qRT-PCR |
| ABFP2584       | TTTGCGGATGGGTTTCCA | primer for qRT-PCR |
| ABFP2660       | ATAGTATGTACAGGACCG | primer for qRT-PCR |
| ABFP2660       | ATGGTCTTACGGATGCAAG | primer for qRT-PCR |
| ABFP1331       | TGCTGAAGACAGAAATGAC | primer for qRT-PCR |
| ABFP1331       | AGTCTTTCGCTTTCTGTG | primer for qRT-PCR |
| AB16S_rRNAFP   | CAGCGGTGCGAGGATGCTG | primer for qRT-PCR |
| AB16S_rRNAFP   | CTTAAGGGCGATGGATCCTT | primer for qRT-PCR |
| FPAB0229       | AGCGATATAGCCAAGTGG | primer for qRT-PCR |
| FPAB0229       | CCTTGCAGACCCAAAGATC | primer for qRT-PCR |
| FPAB1505       | TCAATGGCAACAACACCCTT | primer for qRT-PCR |
| FPAB1505       | AAGTCCAGGACCCGACATC | primer for qRT-PCR |
| FPAB3401       | TGCAACCCCAGAATAATAA | primer for qRT-PCR |
| FPAB3401       | CTGGCGTCCATGCTGTTAT | primer for qRT-PCR |
| FPAB3342       | AAGAGAAACACCGGTGCT | primer for qRT-PCR |
| FPAB3342       | TCAAGGCGCCAAAGATACC | primer for qRT-PCR |
| FPAB1627       | TCTTGCAGGTCGCTGGG | primer for qRT-PCR |
| FPAB1627       | TGATTTTTCAGGGGT | primer for qRT-PCR |
| FPpWHorI       | TACGGTTTGCATGCAAGATATCTAT | Primer for Ori cloning |
| RPpWHorI       | TAATGGTATGGTTAACTTGCTG | Primer for Ori cloning |
| AbsR25FP       | TTTTGGTCGACCATAGTGGTTATATATTACGGGAGTTTACTTTCCTG | Primer for AbsR25 |
| AbsR25R     | TGCTAAGGCTGGAAATAAAATACCAAGAAAAAGGCTAC | Primer for promoter |
| pBADFP         | ATATGAGCTCATGCAATGTTGCTCGTCAATG | Primer for promoter |
| pBADRP         | ATATGCATGAGGCAAAATACCAAGAAAAAGGCTAC | Primer for promoter |
| AbsR11_probe   | ATGCCCTTAAACAGTACGCGGTCACTCATGAGAG-TCTAATTTTTTAACTAGCTTCTCACATATCGC | Sequence for synthesis of riboprobe for AbsR11 sRNA detection |
| AbsR25_probe   | TCTTTTTTAAATCTATGGAATTGACCGATGCTC | Sequence for synthesis of riboprobe for AbsR25 sRNA detection |
| AbsR28_probe   | AAAAGAGGAGACATCAGCCAAAACACAGTAAACACAC-AGTAGAAGGTCGTCAGCTCGTGC | Sequence for synthesis of riboprobe for AbsR28 sRNA detection |

doi:10.1371/journal.pone.0093833.t001
Cincinnati, Ohio, USA) according to manufacturer’s instructions, followed by DNase I (Fermentas, USA) treatment for removal of contaminating genomic DNA. All other steps of qRT-PCR were performed as mentioned earlier.

Results

Prediction of putative sRNAs using bioinformatic approach

There are several experimental strategies used for prediction of sRNA that are based on shotgun cloning and microarray methods but computational predictions and validation with Northern blotting have been popularly used for identification of sRNAs in many bacteria [61]. The basis for genome wide screening of small RNAs is their sequence or structural conservation among closely related species. The algorithms that are based on sequence conservation like sRNAPredict [50,55] and GMMI [62] first search the IGRs (intergenic regions) for transcriptional signal sequences like promoters and terminators and then predict nucleotide conservation. Since the sequence homology is typically based on the primary structure of RNA, so the precision of this method may not be sufficient [63]. Hence some algorithms like QRNA are required [64] that search for phylogenetic conservation of secondary structure. In this study, we employed a combination of sRNAPredict and QRNA algorithms to increase the prediction accuracy.

When this study for sRNA prediction was initiated, among Acinetobacter species, genome of only Acinetobacter baumannii ATCC 17978 was sequenced. Therefore closely related bacterial genomes were taken for comparative analysis on the basis of 16S rRNA gene sequences. In the meantime, other Acinetobacter species genome sequences also became available. There were, thus, two parallel studies for sRNA prediction by using the same methodology (using QRNA and sRNAPredict program), one with closely related bacterial genomes and another with Acinetobacter spp. (Acinetobacter baylyi ADP1 and Acinetobacter sp. DR1). Overlapping results from the two studies increased the level of confidence in the prediction of putative sRNAs and narrowed down the range of putative sRNAs to be verified.

In the study reported here, several steps were employed for prediction of sRNAs of A. baumannii (Figure 1). The phylogenetic relationship of A. baumannii with other bacteria was determined by aligning their 16S rRNA gene sequences. The alignment showed that A. baumannii was closest to P. aeruginosa, S. maltophilia and B. multivorans (data not given). The 16S rRNA gene sequences of A. baumannii and P. aeruginosa showed 88% similarity (100% query coverage) whereas the rest of the sequences showed 85% similarity (100% query coverage) with A. baumannii ATCC 17978 genome. Initially, comparative genomics was started with these genomes. Similar comparative genomics for sRNA prediction was carried out again when genome sequences of A. baylyi ADP1 and Acinetobacter sp. DR1 became available.

IGRExtract was used for extracting intergenic regions (IGRs) from respective genomes for producing IGR databases. These databases were used as subject sequences in BLAST search and converted into coordinates of conserved sequences in the chromosome by sRNAPredict program. IGRs of A. baumannii were extracted and used as query for BLAST search against databases of P. aeruginosa, B. multivorans and S. maltophilia genomes.

Figure 1. Schematic representation of bioinformatic approach used for small RNA species prediction in Acinetobacter baumannii ATCC 17978. The first step was to determine the inter-genic regions (IGRs) from the annotated ORFs (Open Reading frames) and then running them through BLAST against genomes of related species. The conserved sequences were subjected to QRNA program wherein conserved secondary structure regions among the pairwise aligned sequences were determined.

doi:10.1371/journal.pone.0093833.g001
BLAST alignments were used as input for QRNA. QRNA classified a total of 99 candidate structural RNA loci in A. baumannii (Table S1). Thus, there are 99 sequences in IGR of A. baumannii with secondary structure conserved across all the three genomes. For further analysis through sRNAPredict3, rNAMotif and TransTermHP were used for predicting rho-independent terminators, and BLAST was done again with the IGR sequences of A. baumannii as database. sRNAPredict gave 164 coordinate pairs (Table S2) of which only seven lie in or have regions predicted by QRNA to have conserved secondary structure. These results were used later for comparison with results from further analysis, done using other Acinetobacters spp. namely Acinetobacter baysii ADP1 and Acinetobacter spp. DR1. The genome sequences of these were taken as a database for BLAST against A. baumannii IGR sequences and provided too many results with the parameters used previously as they were calibrated for 65–100% similar sequences; so default parameters were used which provide 95–100% similar sequences. Running the result through QRNA provided 806 coordinate pairs for conserved secondary structure (Table S3). For sRNAPredict, BLAST was done again with IGR sequences as database and the genomes (A. baysii ADP1 and Acinetobacter spp. DR1) as a query. The result of sRNAPredict was 748 putative sRNAs (Table S4) of which 143 were predicted by QRNA as having conserved secondary structure.

Comparing these 748 with the 164 obtained before gave 149 coordinate pairs with either the same start coordinate or same end coordinate or both. Of these 149, only 31 are such which were predicted by QRNA as having conserved secondary structure (Table 2). On the basis of these programs (sRNAPredict and QRNA), 31 putative sRNA genes were predicted in the genome of A. baumannii ATCC 17978 (Table 2). These 31 putative sRNAs were predicted based on their location in the intergenic region, conserved secondary structure and termination by rho-independent terminator. The genes encoding the sRNA are denoted as absR (1 to 31) for Acinetobacter baumannii small RNA.

Experimental validation of predicted sRNA

The putative sRNAs predicted using sequence data of A. baumannii ATCC 17978 were verified in A. baumannii MTCC 1425 (ATCC 15308) which is an MDR strain. The expression of RNA species was examined in cells grown to lag, early exponential, exponential and stationary phase. The cells were also subjected to stress conditions, of which some are reminiscent of environmental conditions that A. baumannii cells encounter upon invasion of human wounds. These include temperature and osmotic stress. Out of 31 candidate genes, 10 candidate sRNA genes (absR 9, 11, 13, 14, 16, 17, 22, 24, 25, 28) were tested. These genes were chosen for validation as they were the most conserved (having lowest E values) among intergenic regions transcribing sRNA candidates of other sequenced Acinetobacter spp. and closely related bacteria as predicted by BLAST analysis. 78 nucleotides long specific riboprobes were designed for detection of signal in Northern blotting. For Northern blotting, total RNA of four different growth phases (lag, early exponential, exponential and stationary) was tested. Of the ten putative sRNA candidates analyzed, three (AbsR11, 25 and 28) showed positive signal on Northern blots (Figure 2A, B, C).

The predicted length of AbsR11 sRNA was 351 nts (nucleotides) but in Northern blot a positive signal of 200 nts was obtained which was further confirmed by RACE mapping (Figure S1). AbsR11 sRNA was observed during all phases of bacterial growth with almost the same level of expression (Figure 2A). AbsR28 was predicted to be 205 nts long and is located between AbsR2838 (encoding a hypothetical protein) and AbsR2839 (encoding a hypothetical protein), the experimentally proven length is approximately 180 nts (Figure S1). Its expression was also observed in all the three growth phases tested (Figure 2C).

AbsR25 sRNA showed a differential level of expression with no expression in lag phase, basal expression in early exponential and stationary phase and maximum expression in exponential phase (Figure 2B). The length of AbsR25 sRNA predicted by bioinformatic approaches is 333 nts and the location is between AbsR1321 and AbsR1322 genes, encoding putative hemolysin and dihydrodipicolinate synthetase respectively. Northern blotting and RACE mapping showed that the transcript of AbsR25 sRNA was about 164 nts (Figure S1). The shortening in length might be due to processing of 3’ and 5’ ends of sRNA. Expression of AbsR25 was also tested under different conditions of temperature, osmotic and ethidium bromide stress.

For temperature stress, bacterial cultures were grown at 37°C, 45°C and 50°C for 30 minutes. The level of expression of AbsR25 sRNA was same at 30°C, 37°C and 45°C but no expression was observed at 50°C (Figure 3A)

Osmotic stress to A. baumannii cells was provided by growing cells under different concentrations of NaCl (100, 200, 300, 400 and 500 mM). As the concentration of NaCl increased, the expression of AbsR25 increased. This indicates that this sRNA may be necessary for survival of bacteria under osmotic stress conditions (Figure 3B).

For ethidium bromide (EtBr) stress, bacterial cells were grown in a gradient of EtBr concentration (0 µg/ml, 64 µg/ml, 128 µg/ml, 192 µg/ml and 256 µg/ml respectively). An increase in the concentration of EtBr decreased the level of expression of AbsR25 (Figure 3C). This indicates that this sRNA might have a role in the survival of the bacterium under EtBr stress. Since it is well documented in the literature that EtBr is a common substrate of many efflux pump families, the link between sRNA and efflux pumps was explored and the targets of this sRNA were predicted using bioinformatic approaches.

The expression of other predicted sRNAs was not observed in the tested conditions for the present study. These sRNA genes might be expressed in different growth and physiological conditions specific to the cellular needs.

RACE mapping of validated sRNAs

Northern blots only provide information about the expression level and the approximate size of a transcript, but cannot detect the exact position of the 3’ and 5’ ends of sRNA. For precise detection of transcription start and stop site, rapid amplification of cDNA ends (RACE) analysis was performed. After RACE mapping the size of AbsR11, AbsR25 and AbsR28 were predicted to be 200 nts, 164 nts and 180 nts respectively (Figure S1). Sequences of sRNAs and its location in the genome are shown after BLAST analysis (Figure S1).

Distribution of the identified sRNA genes in other bacteria

To determine whether the sRNA genes identified in this study are present in other microorganisms, the complementary DNA sequence of the sRNA genes was used to perform a BLAST search against the NCBI total sequence database (http://www.ncbi.nlm.nih.gov/Genebank/index.html). The results depicted in Table 3 show that none of the three identified sRNAs display sequence similarity with other bacterial species, indicating that these three sRNAs are novel. The result of AbsR11 sRNA BLAST showed that homologous sequences were found in all sequenced Acinetobacter species, indicating that AbsR11 may be Acinetobacter specific sRNA. Homologous gene sequences of absR25 were found.
| sRNA Up ORF name | Up ORF | dist Up ORF | sRNA start | sRNA end | direction sRNA | length sRNA | dist Dn ORF | Dn ORF | Dn ORF name | BLAST expect | Term Type |
|-----------------|--------|-------------|------------|----------|---------------|------------|------------|--------|-------------|--------------|-----------|
| AbR-50S ribosomal protein 1 | >>>> | 0 | 304073 | 304175 | >>>> | 102 | 390 | >>>> | 50S ribosomal protein | 2.00E-155 | R |
| AbR-50S ribosomal protein 2 | >>>> | 1518 | 1439995 | 1440255 | >>>> | 260 | 295 | >>>> | phage putative head morphogenesis protein | 1.40E-120 | R |
| AbR-amino acid ABC transporter permease 3 | <<< | 0 | 2665657 | 2665731 | >>>> | 274 | 11 | <<< | lysine/arginine/ornithine ABC transporter periplasmic-binding protein | 5.50E-120 | R |
| AbR-lysine-specific permease 4 | >>>> | 36 | 3285829 | 3285889 | >>>> | 60 | 280 | >>>> | hypothetical protein | 7.50E-15 | R |
| AbR-hypothetical protein 5 | <<< | 0 | 3765974 | 3766394 | >>>> | 274 | 11 | <<< | putative pirin-like protein | 5.40E-157 | R |
| AbR-uracil transport protein 6 | >>>> | 8 | 3965175 | 3965381 | <<< | 206 | 113 | >>>> | putative transcriptional regulator | 4.50E-53 | R |
| AbR-glucosamine–fructose-6-phosphate aminotransferase 7 | >>>> | 608 | 3904650 | 3904740 | <<< | 90 | 2343 | >>>> | hypothetical protein | 5.80E-200 | R |
| AbR-hypothetical protein 8 | <<< | 392 | 3766366 | 3766439 | <<< | 73 | 0 | <<< | putative pirin-like protein | 5.40E-157 | R |
| AbR-putative sensory transduction histidine kinase 9 | <<< | 1559 | 3362935 | 3363158 | <<< | 223 | 161 | >>>> | hypothetical protein | 8.30E-293 | R |
| AbR-transcriptional factor 10 | >>>> | 17 | 2543367 | 2543541 | <<< | 174 | 195 | <<< | tRNA uridine 5-carboxymethylaminomethyl modification enzyme GidA | 5.40E-62 | R |
| AbR-hypothetical protein 11 | >>>> | 35 | 2529183 | 2529534 | <<< | 351 | 0 | >>>> | cytochrome c ubiquinol oxidase subunit II | 1.80E-202 | R |
| AbR-hypothetical protein 12 | >>>> | 1307 | 1956579 | 1956712 | <<< | 133 | 232 | >>>> | putative methyltransferase | 4.40E-150 | R |
| AbR-OmpA/MotB 13 | >>>> | 33 | 1398936 | 1399168 | <<< | 232 | 75 | >>>> | hypothetical protein | 9.60E-34 | R |
| AbR-hypothetical protein 14 | <<< | 1151 | 940263 | 940781 | <<< | 518 | 12 | <<< | putative transcriptional regulator | 1.90E-16 | R |
| AbR-putative transposase 15 | >>>> | 30 | 798366 | 798883 | <<< | 517 | 16 | <<< | dihydropteroate synthase | 4.00E-240 | R |
| AbR-acyl-CoA synthetase 16 | <<< | 156 | 639378 | 639744 | <<< | 366 | 401 | >>>> | threonyl-tRNA synthetase | 3.60E-233 | R |
| AbR-OGF1 ATP synthase subunit epsilon 17 | >>>> | 20 | 178139 | 178326 | <<< | 187 | 0 | >>>> | hypothetical protein | 1.10E-106 | R |
| AbR-putative phosphatase 18 | <<< | 403 | 501115 | 501198 | <<< | 83 | 2 | <<< | putative alkaline phosphatase | 4.70E-130 | T |
| AbR-integration host factor subunit alpha 19 | >>>> | 2356 | 654179 | 654249 | <<< | 70 | 0 | <<< | hypothetical protein | 0 | T |
| AbR-Transposase 20 | >>>> | 312 | 800944 | 801007 | <<< | 63 | 1 | <<< | rRNA | 1.20E-42 | T |
| sRNA Up ORF name | Up ORF dir. | dist Up ORF | sRNA start | sRNA end | sRNA direction | length sRNA | dist Dn ORF | Dn ORF dir. | Dn ORF name | BLAST expect | Term Type |
|-----------------|-------------|-------------|-------------|----------|---------------|-------------|-------------|-------------|-------------|--------------|-----------|
| AbsR-21 putative homocysteine 5'-methyltransferase family protein | <<<< | 862 | 10718391071925 | <<<< | 86 | <<<< | malate:quinone oxidoreductase | 6.40E-221 | T |
| AbsR-22 major facilitator superfamily fosmidomycin/multidrug transport protein | >>>> | 1084 | 11150251115571 | >>>> | 546 | 26 | mutarotase precursor | 6.40E-20 | B |
| AbsR-23 OmpA/MotB | >>>> | 10 | 13989131399168 | <<<< | 255 | 75 | hypothetical protein | 9.60E-34 | T |
| AbsR-24 putative hemolysin | >>>> | 0 | 15488161548878 | >>>> | 62 | 910 | dihydrodipicolinate synthetase | 0 | B |
| AbsR-25 putative hemolysin | >>>> | 0 | 15488161549149 | >>>> | 333 | 639 | dihydrodipicolinate synthetase | 0 | B |
| AbsR-26 hypothetical protein | >>>> | 1289 | 19565611956712 | <<<< | 151 | 232 | putative methyltransferase | 4.40E-15 | T |
| AbsR-27 hypothetical protein | >>>> | 35 | 25291832529534 | <<<< | 351 | 0 | cytochrome c ubiquinol oxidase subunit II | 1.80E-202 | R |
| AbsR-28 lysine-specific permease | >>>> | 36 | 32858293286033 | >>>> | 204 | 136 | hypothetical protein | 7.50E-15 | B |
| AbsR-29 hypothetical protein | >>>> | 14 | 34631913463452 | >>>> | 261 | 169 | putative vanillate O-demethylase oxygenase subunit | 1.50E-12 | T |
| AbsR-30 S-methylmethionine APC transporter | <<<< | 404 | 36235313623594 | >>>> | 63 | 12 | hypothetical protein | 3.90E-09 | T |
| AbsR-31 S-methylmethionine APC transporter | <<<< | 397 | 36235243623606 | <<<< | 82 | 0 | hypothetical protein | 1.80E-15 | T |

Up ORF: Upstream Open Reading frame; dist Up ORF: distance of sRNA from Upstream Open Reading frame; UpORF dir: Upstream Open Reading frame direction; dir sRNA: direction of small RNA expression; len sRNA: length of sRNA; dist Dn ORF: distance of sRNA from downstream open reading frame.
only in A. baumannii strains except SDF strain, indicating that AbsR25 could be species specific sRNA. In case of AbsR28, homologous sequences were found in all sequenced Acinetobacter species except Acinetobacter sp. ADP1 suggesting that this could also be Acinetobacter specific sRNA.

Prediction of AbsR25 sRNA target genes

In silico analysis was performed for prediction of AbsR25 target genes. For more specific results, two different programs (sRNA-TargetNB and IntaRNA) were used. sRNA-TargetNB uses Naïve Bayes probabilistic method for sRNA target gene prediction and takes secondary structure profile as the feature [59]. IntaRNA predicts interactions between two RNA molecules, and the scoring is based on the hybridization free energy and accessibility of the interaction sites in both molecules [60]. By using these two target prediction tools, 62 common putative target genes were predicted for AbsR25 (Table S5). The combined in silico data constitute the hypothetical regulon for AbsR25 and these genes were experimentally verified.

Confirmation of AbsR25 target genes

It is well characterized that expression of target genes is controlled by base pairing of sRNA with the cognate target mRNA. Base pairing of sRNA with its target mRNA not only affects translation but also leads to change in the stability of both sRNA and mRNA. To experimentally identify potential target genes of AbsR25 sRNA, the transcriptomes of A. baumannii were compared. As the levels of AbsR25 were found to decrease with increasing concentration of EtBr (in the earlier experiment described above, Figure 3C), the cells were grown in nutrient broth, in the presence of different concentrations of EtBr and in the absence of EtBr. For expression analysis of putative target genes, qRT-PCR was used. Out of 62 putative target genes, initially 11 genes were screened on the basis of having higher hybridization energy predicted by the IntaRNA program (Table 4). Out of these mRNAs, nine (AIS_0229, AIS_1331, AIS_3401, AIS_1627, AIS_2269, AIS_2660, AIS_3251, AIS_3342, AIS_1791) showed increased expression (Figure 4) of mRNA transcript relative to control (sample prepared without EtBr treated culture) (Table 4). The maximum change in expression (27.09 fold increase) was observed in case of AIS_0229 (hypothetical protein), 7.67 fold increase in expression in case of AIS_1331 (AIS_1627, AIS_2269, AIS_2660, AIS_3251, AIS_3342) showed increased expression and AIS_1791 (hypothetical protein) gene. Although the change in expression was observed in many genes, consistent increase in various concentrations of EtBr was observed only in case of AIS_1331 (major facilitator superfamily transporter gene) (Figure 4). The expression of AbsR25 sRNA was also tested in the same conditions which showed a decrease in its expression as the concentration of EtBr was increased from 0 μg/ml to 256 μg/ml. Conversely, as the concentration of EtBr was increased, the expression of AIS_1331 gene which encodes a major facilitator superfamily transporter or efflux pump increased at least six fold. This shows that AbsR25 negatively regulates the efflux pump transporter, AIS_1331 (major facilitator superfamily transporter) which controls the efflux of EtBr.
Since EtBr itself can affect the expression of some genes, therefore AbsR25 was over-expressed to determine bona-fide interactions with the target genes. A. baumannii strain designed to express higher amounts of AbsR25 was created by directional cloning of absR25 gene in pPROBE-PBAD. Depending upon the cloning strategy, expression of absR25 was controlled by P_{BAD}.

Table 3. Distribution and conservation of the identified sRNA gene sequence (absR25) in closely related bacteria.

| Bacterial strains | AbsR11 (Identity out of 100% query coverage) | AbsR25 (Identity out of 100% query coverage) | AbsR28 (Identity out of 98% query coverage) |
|-------------------|---------------------------------------------|---------------------------------------------|---------------------------------------------|
| Aba 17978         | 100%                                        | 98%                                         | 98%                                         |
| Aba AYE           | 99%                                         | 98%                                         | 99%                                         |
| Aba ACICU         | 100%                                        | 98%                                         | 99%                                         |
| Aba AB0057        | 99%                                         | 98%                                         | 99%                                         |
| Aba AB307-0294    | 99%                                         | 98%                                         | 99%                                         |
| Aba 1656-2        | 100%                                        | 98%                                         | 99%                                         |
| Aba MDR-ZJ06      | 100%                                        | 98%                                         | 99%                                         |
| Aba TCDC-AB0715   | 100%                                        | 98%                                         | 99%                                         |
| Aba MDR-TJ        | 100%                                        | 98%                                         | 99%                                         |
| Aba SDF           | 99%                                         | N                                           | 98%                                         |
| Aca PHEA-2        | 96%                                         | N                                           | 89%                                         |
| Aol DRI           | 95%                                         | N                                           | 92%                                         |
| A sp. ADPI        | 84%                                         | N                                           | N                                           |

Here in table; Aba, Acinetobacter baumannii; Aca, Acinetobacter calcoaceticus; Aol, Acinetobacter oleivorans; N, no similar sequence found.

doi:10.1371/journal.pone.0093833.t003
Selected target genes for expression analysis by qRT-PCR.

Normalized relative expression of target mRNAs & AbsR25 in the presence of EtBr concentrations (µg/ml) 0 64 128 192 256 Uninduced Induced

| S. No. | Putative target mRNA | EtBr concentrations (µg/ml) | 0 | 64 | 128 | 256 | 192 | 256 |
|--------|----------------------|-----------------------------|---|----|-----|-----|-----|-----|
| 1      | A1S_3401 (hypothetical protein) | 1 | 1.27 | 6.2 | 0.1 | 0.06 | 1 | 0.14 |
| 2      | A1S_3251 (amino acid transporter LysE) | 1 | 1.51 | 5.54 | 0.16 | 0.15 | 1 | 0.46 |
| 3      | A1S_1791 (putative tartrate symporter MFS superfamily protein) | 1 | 1.84 | 0.66 | 0.8 | 0.38 | 1 | 0.86 |
| 4      | A1S_2269 (RtcR - Regulator of RNA terminal phosphate cyclase) | 1 | 1.87 | 0.86 | 1.53 | 1.59 | 1 | 1.23 |
| 5      | A1S_1505 (hypothetical protein) | 1 | 1.04 | 0.32 | 0.69 | 0.56 | 1 | 0.86 |
| 6      | A1S_0229 (hypothetical protein) | 1 | 1.21 | 0.52 | 0.33 | 1.63 | 1 | 0.86 |
| 7      | A1S_3342 (putative arsenate reductase) | 1 | 1.21 | 3.34 | 3.34 | 0.58 | 0.91 | 1 | 0.86 |
| 8      | A1S_1627 (hypothetical protein) | 1 | 1.21 | 0.52 | 0.33 | 0.07 | 0.06 | 1 | 0.86 |
| 9      | A1S_2660 (RND efflux transporter) | 1 | 1.21 | 0.52 | 0.33 | 0.07 | 0.06 | 1 | 0.86 |
| 10     | A1S_1331 (major facilitator superfamily transporter) | 1 | 1.21 | 0.52 | 0.33 | 0.07 | 0.06 | 1 | 0.86 |
| 11     | AbsR25 sRNA | 1 | 0.08 | 0.25 | 0.07 | 0.07 | 0.07 | 1 | 0.86 |

Discussion

In the present study we report the integration of computational and experimental methods to identify novel sRNAs in the nosocomial pathogen; A. baumannii. The bioinformatic prediction of sRNAs was based on screening the intergenic regions (IGR) of A. baumannii ATCC 17978 genome for sequence conservation, rho-independent termination and conserved secondary structures. As a measure to enhance the accuracy, two different query sets for prediction were prepared. First set contained the sequences from species closely related to A. baumannii (P. aeruginosa, B. multivorans and S. maltophilia) while the other set contained sequences from two A. baumannii strains (A. baylyi ADP1 and Acinetobacter spp. DR1).

Prediction was carried out on both the sets and only the common results obtained from both the sets were considered for further analysis. Two web based programs (sRNApredict and QRNA) were used for increasing the sensitivity of the sRNA prediction. The sRNApredict algorithm is based on two types of information, first the location of transcriptional signal and second primary sequence conservation of IGRs. On the other hand QRNA algorithm is based on conservation of secondary structures of RNA for prediction. As a result of this exercise a list of 31 putative sRNA molecules was generated which satisfied the set conditions and contained common sequences predicted as sRNA from both the groups.

On the basis of sequence conservation and lower E-value in BLAST analysis, 10 out of the 31 putative sRNA molecules were selected for experimental validation. Northern blotting was performed to detect the expression of the selected 10 putative sRNAs in A. baumannii MTCC 1425 (ATCC 15308). The MTCC 1425 strain is an MDR strain showing resistance to 12 antibiotics belonging to different chemical classes (data not shown). Positive signal on Northern blot was detected for three putative sRNA species namely; AbsR11, AbsR25 and AbsR26. The sizes of all the three molecules were shorter than the predicted size which might be due to processing at the 5’- or the 3’- end. This size difference...
was confirmed by RACE mapping. The size of AbsR11, AbsR25 and AbsR28 was found to be 200 nts, 164 nts and 180 nts, respectively corresponding to the sizes predicted by Northern blot. BLAST analysis revealed that AbsR11 and AbsR28 are conserved among the *Acinetobacter* spp. but no such homology existed in other genera, making them genus specific. However, AbsR25 was found conserved only among *A. baumannii* strains making this sRNA species specific (Table 3). Since no similarity of these sRNAs with already identified sRNAs was found in other bacteria, these sRNAs are novel in nature.

**Figure 4. Real time expression of selected target genes.** Relative expression of *in silico* predicted putative target genes [A1S_3401 (hypothetical protein), A1S_3251 (amino acid transporter LysE), A1S_1791 (putative tartrate symporter MFS superfamily protein), A1S_2269 (RtcR - Regulator of RNA terminal phosphate cyclase), A1S_1505 (hypothetical protein), A1S_0229 (hypothetical protein), A1S_3342 (putative arsenate reductase), A1S_1627 (hypothetical protein), A1S_2584 (major facilitator superfamily family drug transporter), A1S_2660 (RND efflux transporter), A1S_1331 (major facilitator superfamily transporter)] of AbsR25 sRNA under different concentrations of EtBr.

doi:10.1371/journal.pone.0093833.g004
The expression of AbsR11 and AbsR28 was independent of the stage of growth of *A. baumannii* as the Northern blot using RNA from different growth phases (lag, early exponential, exponential and stationary) showed almost equal level of expression. However, in case of AbsR25, the expression of the said sRNA was not detectable in lag phase, basal in early exponential and stationary phase and maximum in exponential phase. This intriguing observation about AbsR25 led to further investigation on the molecule. The differential expression may indicate its role in stress conditions when there is a scarcity of nutrients. To establish the role of AbsR25 in stress response, its expression profile was studied under gradients of different stress conditions like osmotic, temperature and EtBr stress. These conditions represent the stress that the pathogen is likely to encounter when it makes entry into the host. No change in the expression of AbsR25 was observed in the temperature range of 30°C to 45°C. At 50°C expression of AbsR25 could not be detected which may be due to the difficulty of the bacterial strain to grow (Figure 3A). In contrast, the expression of AbsR25 varied with the concentration of salt (NaCl) in the medium. An increase in the expression of AbsR25 was seen with progressively increasing concentrations of NaCl. This further strengthens the hypothesis that the small RNA may be involved in the cellular mechanism to cope up with environmental stress.

A significant observation was made in the case of EtBr stress. The expression of AbsR25 was found inversely related to the concentration of EtBr in the growth medium of *A. baumannii* (Figure 3C). This observation suggests that AbsR25 might play an important role in efflux pump regulation and drug resistance. Interestingly, most of the predicted targets for AbsR25 were efflux pumps with high hybridization energy. This indicates that AbsR25 might be regulating, directly or indirectly, the expression of efflux pumps. To validate this hypothesis, qRT-PCR was performed with the cDNA of the EtBr stressed culture. Out of the mRNA target genes tested, maximum variation in expression was shown by AIS_0229 (hypothetical protein), AIS_1331 (major facilitator superfamily transporter), AIS_3401 (hypothetical protein) and AIS_3251 (amino acid transporter LysE). Further experiments will be needed to verify these targets. qRT-PCR results reveal that the presence of EtBr in culture medium of *A. baumannii* resulted in decreased AbsR25 expression level and upregulated AIS_1331 mRNA levels which specifies an MFS pump in *A. baumannii*. It is evident that the expression of mRNA specifying AIS_1331 transporter increased uniformly with the increase in concentration of EtBr which also coincided with a steady decrease in the expression of AbsR25 sRNA. It is proposed that this sRNA might be involved in negative regulation of the AIS_1331 transporter. In the absence of EtBr in the culture, sRNA expression was found high enough to prevent the expression of the transporter, however, when the concentration of EtBr increased, the expression of sRNA decreased and the transporter is available for fluxing out the EtBr for survival of the bacterium.

However, EtBr is known to affect the expression of a lot of genes. Therefore, to overcome the effect of EtBr itself on the expression of said genes, AbsR25 was overexpressed in *A. baumannii* and the effect of its overexpression was studied on target genes. The change in expression of the target genes was in accordance with the observations made earlier (Table 4). The increased expression of AbsR25 resulted in 9, 7.14, 5, 2.3, 1.9 and 1.1 fold repression of AIS_0229, AIS_3401, AIS_3342, AIS_1331, AIS_2660, AIS_1791 genes respectively. As is evident from the experiment, the affected genes include five putative transporters belonging to different families. Out of these putative transporters, AIS_1331 has been confirmed as an MFS transporter in a separate study in our group (unpublished results). On the basis of these observations, AbsR25 could be implicated in negative regulation of the expression of AIS_1331 transporter gene, which is in close agreement with EtBr treatment of *A. baumannii* cells. The repression of the genes indicates that the regulation of AIS_1331 transporter gene by AbsR25 sRNA is achieved at the mRNA level. Putative pairings between AbsR25 and AIS_1331 mRNA have been identified but require direct experimental support. AbsR25 might also regulate other targets encoded at separate locations on the chromosome. Since it is well established that efflux pumps regulate clinically relevant resistance to antibiotics [65], these findings provide a new perspective for future research into the mechanisms responsible for antibiotic resistance.
The identification, experimental confirmation, and preliminary investigation of novel and distinctive sRNAs from the nosocomial pathogen *A. baumannii* contribute to the ongoing attempt to characterize and comprehend the function of these significant RNA molecules. This is the first report of novel sRNAs in *A. baumannii* and preliminary result of involvement of a sRNA, AbsR25, in regulating an efflux pump in clinically significant MDR strain. Further experiments are needed to disclose elucidations to the exciting observation of the involvement of AbsR25 sRNA in regulation of expression of efflux pumps and hence in multiple drug resistance.

**Supporting Information**

Figure S1 Sequences of sRNAs and its location in the genome after BLAST analysis. (DOC)

Table S1 QRNA output for first study. (XLSX)

Table S2 Output of sRNAPredict for first study. (XLSX)

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Table S3 QRNA output file for second study. (XLSX)

Table S4 sRNAPredict output for second study. (XLSX)

Table S5 Putative target genes of AbsR25 predicted by bioinformatic analysis. (XLSX)

Table S6 BLAST analysis of six repressed target genes. (DOC)

Acknowledgments

We are thankful to Professor Steven E. Lindow for the kind gift of pPROBE-NT plasmid.

**Author Contributions**

Conceived and designed the experiments: RP NKN RS SA. Performed the experiments: RS SA AS SDP. Analyzed the data: RP NKN RS AS PKJ. Contributed reagents/materials/analysis tools: RP NKN RS PKJ SA. Wrote the paper: RP SDP NKN.
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