# Identification of antibiotic resistance genes in the multidrugresistant *Acinetobacter baumannii* strain, MDR-SHH02, using whole-genome sequencing

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**Abstract.** This study aimed to investigate antibiotic resistance genes in the multidrug-resistant (MDR) Acinetobacter baumannii (A. baumanii) strain, MDR-SHH02, using whole-genome sequencing (WGS). The antibiotic resistance of MDR-SHH02 isolated from a patient with breast cancer to 19 types of antibiotics was determined using the Kirby-Bauer method. WGS of MDR-SHH02 was then performed. Following quality control and transcriptome assembly, functional annotation of genes was conducted, and the phylogenetic tree of MDR-SHH02, along with another 5 A. baumanii species and 2 Acinetobacter species, was constructed using PHYLIP 3.695 and FigTree v1.4.2. Furthermore, pathogenicity islands (PAIs) were predicted by the pathogenicity island database. Potential antibiotic resistance genes in MDR-SHH02 were predicted based on the information in the Antibiotic Resistance Genes Database (ARDB). MDR-SHH02 was found to be resistant to all of the tested antibiotics. The total draft genome length of MDR-SHH02 was 4,003,808 bp. There were 74.25% of coding sequences to be annotated into 21 of the Clusters of Orthologous Groups (COGs) of protein terms, such as 'transcription' and 'amino acid transport and metabolism'. Furthermore, there were 45 PAIs homologous to the sequence MDRSHH02000806. Additionally, a total of 12 gene sequences in MDR-SHH02 were highly similar to the sequences of antibiotic resistance genes in ARDB, including genes encoding aminoglycoside-modifying enzymes [e.g., aac(3)-Ia, ant(2'')-Ia, aph33ib and aph(3')-Ia],

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β-lactamase genes (*bl2b\_tem* and *bl2b\_tem1*), sulfonamideresistant dihydropteroate synthase genes (*sul1 and sul2*), *catb3* and *tetb*. These results suggest that numerous genes mediate resistance to various antibiotics in MDR-SHH02, and provide a clinical guidance for the personalized therapy of *A. baumannii*-infected patients.

#### Introduction

Acinetobacter baumannii (A. baumanii) is a notable pathogen that causes hospital-acquired infections among immune-compromised patients, accounting for 5% of Gram-negative infections (1). Due to a strong resistance to desiccation and multiple antibacterial agents, the widespread dissemination of multidrug-resistant (MDR) A. baumanii strains has been a threat to hospitalized patients in recent years (2).

Antibiotic resistance determinants play pivotal roles in whether or not A. baumannii will flourish in the host (3). For instance, the expression of  $\beta$ -lactamase genes [e.g., oxacillinase (OXA)-235 gene,  $bla_{\text{OXA-51}}$  and  $bla_{\text{TEM-1}}$ ] has been shown to be involved in antibiotic resistance (4-6). An 86-kb region AbaR resistant to heavy metal and antibiotics has been found in a MDR isolate AYE (7), indicating the important role of AbaR in the spread of A. baumannii in hospitals (8,9). Furthermore, other resistance determinants, such as macrolide (msrA/msrB), aminoglycoside (e.g., aacCl, armA, and aphAl) and tetracycline [e.g., tet (39), tet (A), and tet (B)] have been identified in various A. baumannii isolates (10).

Bacterial whole-genome sequencing (WGS) has enhanced our ability to evaluate antibiotic resistance determinants. For example, a AbaR-type genomic resistance island, AbaR22, has been identified in the MDR *A. baumannii* strain, MDR-ZJ06, via WGS (11). A whole-genome comparison detected 18 putative single nucleotide polymorphisms (SNPs) between 2 pre- and post-therapy *A. baumannii* isolates (12). Furthermore, 10 types of AbaR resistance islands were identified in 2 *A. baumannii* isolates using WGS (13). Despite increased research on antibiotic resistance determinants in *A. baumannii*, the molecular mechanisms of antibiotic resistance in MDR *A. baumannii* are

not yet fully understood, and various antibiotic resistance genes have not been detected.

In the present study, we applied WGS to obtain the whole genomic sequence of the MDR *A. baumannii* strain, MDR-SHH02, isolated from a patient with breast cancer. Furthermore, the antibiotic resistance of MDR-SHH02 to multiple antibiotics was determined, and potential antibiotic resistance genes in MDR-SHH02 were predicted. The results of our study may enhance our understanding of the molecular mechanisms of antibiotic resistance in MDR *A. baumannii*, and provide a clinical guidance for the therapy of *A. baumannii*-infected patients.

## Materials and methods

Isolation and identification of A. baumanii strain. The clinical MDR A. baumanii strain, named MDR-SHH02, was isolated from the blood obtained from a 65-year-old woman with terminal-stage breast cancer at Shanghai Sixth People's Hospital Affiliated to Shanghai Jiaotong University, Shanghai, China. This patient had received a double mastectomy and nearby lymph node excision. After being discharged from the hospital, this patient was hospitalized again due to symptoms of fever, cough (lasting for days) and shortness of breath. During her second hospital administration, she received several antimicrobial therapies, including maxipime, impenem, methylprednisolone, levofloxacin and cefoperazone-sulbactam sodium. A. baumanii was positive in the blood culture and sputum culture. The results of the antimicrobial susceptibility test revealed that the A. baumanii strain was resistant to multiple commonly used antibiotics. This study was approved by the Shanghai Health and Family Planning Commission Foundation (Shanghai, China), and informed consent was obtained from the patient.

The isolated strain was inoculated onto blood agar plates and then incubated in an atmosphere of 5% CO<sub>2</sub> at 35°C for 48 h. Afterwards, this strain was identified using morphological and biochemical tests according to standard methods (14). Colonies with typical morphological and biochemical characteristics of *Acinetobacter* were cultivated on 5% sheep blood agar and identified using an automated Microscan® system (Dade Behring, Inc., West Sacramento, CA, USA). The *A. baumanii* strain was stored at -70°C in skim milk for further analyses.

Antibiotic resistance test for A. baumanii MDR-SHH02. According to the Clinical and Laboratory Standards Institute (CLSI) guidelines (15), disc diffusion assay (DDA) with dry wafers saturated by 19 types of antibiotics, including gentamicin (10 µg/wafer), tobramycin (30 µg/wafer), amikacin (30 µg/wafer), ampicillin-sulbactam (10/10 µg/ wafer), ceftazidime (30  $\mu$ g/wafer), ciprofloxacin (5  $\mu$ g/wafer), levofloxacin (5 µg/wafer), imipenem (10 µg/wafer), meropenem (10 µg/wafer), piperacillin/tazobactam (100/10 µg/wafer), ticarcillin/clavulanic acid (75/10 µg/wafer), cefepime (30 µg/wafer), cefotaxime (30 µg/wafer), ceftriaxone (30 µg/wafer), doxycycline (30 µg/wafer), minocyline (30 µg/wafer), tetracycline (30μg/wafer),piperacillin(100μg/wafer)andtrimethoprim-sulfamethoxazole (1.25/23.75 µg/wafer) (all from Oxoid, Ltd., Basingstoke, UK), were carried out using the Kirby-Bauer (KB) method, as previously described (16). Briefly, Mueller-Hinton agar (Oxoid, Ltd.) plates were overlaid with the inocula of the clinical *A. baumanii* strain, and the turbidity of the inocula was equivalent to the 0.5 McFarland standard. Subsequently, dry wafers saturated by antibiotics were placed on the surface of the agar, and plates were placed in an atmosphere of 5% CO<sub>2</sub> at 35°C. Following 24 h of culture, the diameter of the inhibition zone around each wafer was measured according to the CLSI criteria (15). In this test, *Escherichia coli* ATCC 25922, ATCC 35218 and *Pseudomonas aeruginosa* ATCC 27853 obtained from the Clinical Laboratory Center of the Ministry of Health were used as reference strains.

DNA preparation, library construction and sequencing. The genomic DNA of A. baumanii MDR-SHH02 was extracted using a bacterial genomic DNA purification kit (Tiangen Biotech Co., Ltd., Beijing, China) according to the manufacturer's instructions. The Illumina sequencing library was then prepared using the Nextera™ DNA Sample Preparation kit (Illumina®-Compatible). Paired-end dual index 2x90 bp sequencing was fulfilled following the Illumina HiSeq 2000. Sequencing was performed by Beijing Genomics Institute (BGI; Shenzhen, China). The sequencing data were uploaded to the public database the National Center for Biotechnology Information (NCBI; http://www.ncbi.nlm.nih.gov/) under the BioProject PRJNA256112 with BioSample accession no. SAMN02991371.

Quality control. For the raw sequencing data, the reads were cleaned by removing the empty reads, adapter sequences and reads with  $n\ge10\%$  using the SeqPrep program (https://github.com/jstjohn/SeqPrep) and Sickle (https://github.com/najoshi/sickle). In addition, the reads were trimmed by discarding the reads containing >30% bases with a Q-value ≤20 in the 3' terminal, as well as reads with adaptor sequences (the length of overlapping sequences between adaptor and read was at least >15 bp, and the number of mismatch bases was <3 bp).

Genome assembly. The clean reads were assembled using the short oligonucleotide analysis package SOAPdenovo (version 2.04; http://soap.genomics.org.cn/). To determine whether the GC content has a significant effect on sequencing randomness or not, the GC content and average depth of the genomic sequence were calculated without repetition as a unit of 500 bp.

Genome annotation. Genes in the assembled genomic sequence were predicted using Glimmer 3.0 (http://www.cbcb.umd.edu/software/glimmer/) (17), which is a system for identifying genes in DNA sequences of microorganism, particularly bacteria, archaea and viruses.

Furthermore,tRNAandrRNA(5S,16S,and23SrRNA)inthe genomic sequence were searched using tRNAscan-SE (http://lowelab.ucsc.edu/tRNAscan-SE/) (18) and RNAmmer 1.2 (http://www.cbs.dtu.dk/services/RNAmmer/) (19), respectively.

Additionally, tandem repeat sequences and clustered regularly interspaced short palindromic repeats (CRISPR) in the genomic sequence were predicted using the Tandem Repeat Finder (http://tandem.bu.edu/trf/trf.html) and CRISPR Finder (http://crispr.u-psud.fr/Server/) software, respectively.

Insertion sequences (ISs) were characterized using the IS Finder database (https://www-is.biotoul.fr//), and the parameter -e was set as 1e-5, identity set as 35%. Besides, protein domains associated with the genomic sequence were predicted using the InterPro database (https://www.ebi.ac.uk/interpro/), and the parameter was set as -appl PfamA.

Functional annotation of genes. Sequence alignment of the amino acid sequences of genes to the Cluster of Orthologous Groups (COGs) of proteins database (http://www.ncbi.nlm.nih.gov/COG/) (20) was performed using the Basic Local Alignment Search Tool (BLASTP; version 2.0) program from NCBI (E-value  $\leq 10-4$ ) (21). We also performed sequence alignment of the amino acid sequences of genes to the NCBI non-redundant (NR) database (E-value  $\leq 10-10$ , identity score  $\geq 35\%$ , and coverage length  $\geq 80\%$ ). If the amino acid sequence of a gene was aligned to multiple sequences in the databases, the optimal result was retained.

Construction of phylogenetic tree. Based on the NCBI 16S rRNA gene database, 16S rRNA gene sequences of 5 A. baumanii strains, including ATCC 17978, ATCC 19606, CIP 70.34, DSM 30007 and A. baumanii JCM 68415, as well as 2 species belonging to Acinetobacter (A. haemolyticus ATCC 17906 and A. bereziniae ATCC 17924), were used to construct the phylogenetic tree, along with MDR-SHH02. Briefly, multiple sequence alignment was performed using ClustalW-2.1 (22). Subsequently, the software package PHYLIP 3.695 (http://evolution.genetics.washington.edu/phylip.html), along with the bootstrap algorithm, was used to construct the maximum likelihood phylogenetic tree, and the phylogenetic tree was visualized by FigTree v1.4.2 (http://tree.bio.ed.ac.uk/software/figtree/).

Prediction of pathogenicity islands (PAIs). The Pathogenicity Island database (PAIDB; http://www.paidb.re.kr/about\_paidb. php), which is a web-based user-friendly resource and widely used for detecting PAIs in newly sequenced genomes (23), was utilized to predict PAIs in the genomic sequence of MDR-SHH02.

Identification of antibiotic resistance genes. To identify potential antibiotic resistance genes in the genomic sequence of MDR-SHH02, sequence alignment of the protein sequences of antibiotic resistance genes in the Antibiotic Resistance Genes database (ARDB; http://ardb.cbcb.umd.edu/) (24) and MDR-SHH02 genomic sequence was conducted using BLASTP (E-value ≤10-10, identity score ≥90%, and coverage length ≥80%).

## Results

Antibiotic-resistance of A. baumanii MDR-SHH02. The antibiotic-resistance assay revealed that the diameter of the inhibition zone for 17 types of antibiotics on MDR-SHH02 plates was 6 mm, apart from levofloxacin and minocyline (diameter, 10 mm) (Table I). According to the CLSI criteria, MDR-SHH02 was resistant to all of the tested antibiotics.

Assembly and annotation of the genomic sequence of A. baumanii MDR-SHH02. During the genome assembly,

Table I. The results of antibiotic-resistance assay for *A. bau-mannii* MDR-SHH02.

Antibiotic name	Diameter of inhibition zone on MDR-SHH02 plate (mm)	Antibiotic-resistance of MDR-SHH02
Ampicillin-sulbactam	6	R
Ceftazidime	6	R
Ciprofloxacin	6	R
Levofloxacin	10	R
Imipenem	6	R
Meropenem	6	R
Gentamicin	6	R
Tobramycin	6	R
Amikacin	6	R
Piperacillin/tazobactam	6	R
Ticarcillin/clavulanic acid	6	R
Cefepime	6	R
Cefotaxime	6	R
Ceftriaxone	6	R
Doxycycline	6	R
Minocyline	10	R
Tetracycline	6	R
Piperacillin	6	R
Trimethoprim- sulfamethoxazole	6	R

Diameter of inhibition zone <15 mm is determined as drug-resistant (R); diameter of inhibition zone between 15 and 17 mm is determined as intermediate (I); and diameter of inhibition zone >17 mm is determined as drug-sensitive (S). A. baumannii, Acinetobacter baumannii.

a total of 85 scaffolds were generated, and the scaffold N50 length was 131,822 bp. The total draft genome length of MDR-SHH02 was 4,003,808 bp, with 38.99% of GC content. There were 3,787 coding sequences, 62 tRNA sequences and 3 rRNA sequences in the genomic sequence. Moreover, 2 CRISPR and 36 tandem repeat sequences, as well as 29 ISs were predicted in the genomic sequence (Table II).

Furthermore, numerous protein domains were predicted in the genomic sequence of MDR-SHH02, such as the AcrB/AcrD/AcrF family (Table III).

Functional annotation of the genomic sequence of A. baumanii MDR-SHH02. According to the COG annotation, 74.25% (2,812/3,787) of coding sequences were annotated into 21 COG terms, which were divided into 3 categories: information storage of processing, cellular processes and signaling, and metabolism. Apart from the category of poorly characterized, most o the coding sequences were annotated into 'transcription' (number of coding sequences, 261) and 'amino acid transport and metabolism' (number of coding

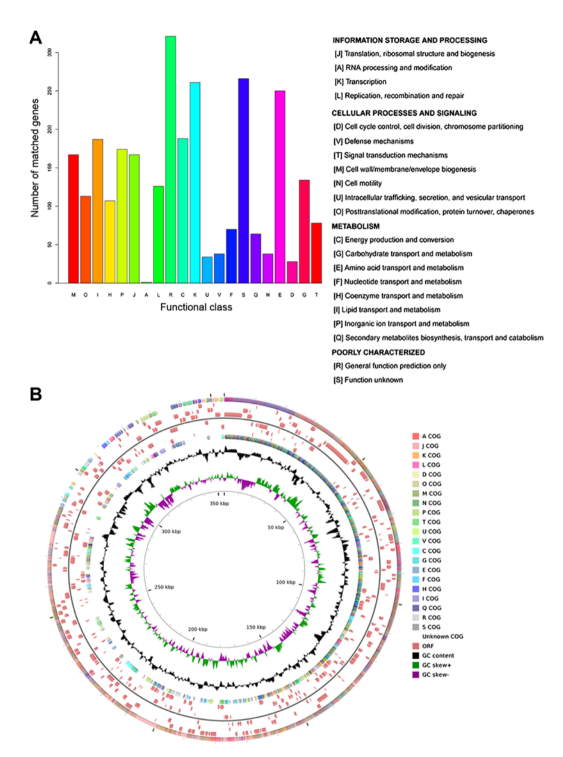


Figure 1. The functional annotation of the genomic sequence of *Acinetobacter baumannii* (*A. baumannii*) MDR-SHH02 using the Cluster of Orthologous Groups (COGs) of proteins database. (A) The bar diagram displaying the COG categories. The x-coordinate represents the COG categories, and y-coordinate represents the number of matched genes; (B) the annotation circle displaying the distribution of COG categories, open reading frames (ORFs) and GC content in the genomic sequence of *A. baumannii* MDR-SHH02. The outermost circle (circle 1) displays the distribution of COG categories on positive strand, and each kind of color represents one COG term, which is annotated as that in (A). Circles 2-4 and 5-7 display the distribution of ORFs on positive and negative strands. Circle 8 displays the distribution of COG categories on negative strand. Circle 9 displays the GC content. Circle 10 displays the GC skew (G-C)/(G+C) (green bars represent positive values, purple bars represent negative values). Circle 11 dislays the the scale in kbp.

sequences, 250) (Fig. 1A). The distribution of COG categories in the genomic sequence of *A. baumannii* MDR-SHH02 is shown in Fig. 1B.

Analysis of the phylogenetic tree. Based on the 16S rRNA gene sequences of A. baumanii in NCBI, the phylogenetic

tree revealed that the genomic sequence of MDR-SHH02 was most similar to the 16S rRNA gene sequence of *A. baumanii* ATCC 17978 (Fig. 2).

Analysis of PAIs. During the process of bacterial infection, PAIs play pivotal roles in the evolution of pathogens and

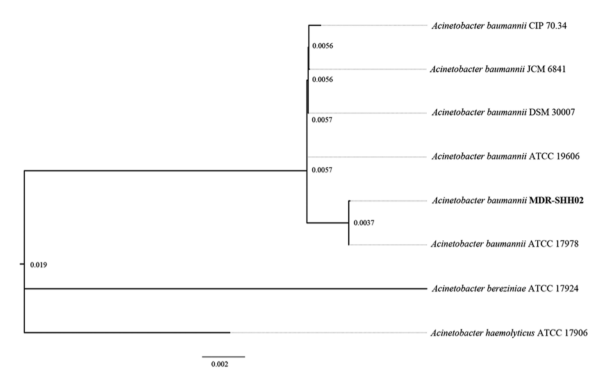


Figure 2. Whole-genome phylogeny of the genomes of *Acinetobacter baumannii* (A. baumannii) MDR-SHH02 and 5 sequenced A. baumannii genomes. The phylogenetic tree was rooted with 2 genomes of *Acinetobacter* species.

Table II. The results of genome assembly and annotation of *A. baumannii* MDR-SHH02.

Feature	Statistics
Length of total draft genome length (bp)	4,003,808
No. of scaffolds	85
Length of scaffold N50 (bp)	131,822
GC content (%)	38.99
No. of coding sequences	3,787
No. of tRNAs	62
No. of rRNAs	3
No. of CRISPR	2
No. of tandem repeat sequences	36
No. of insertion sequences	29

the development of diseases. In the genomic sequence of MDR-SHH02, a total of 45 PAIs homologous to the sequence MDRSHH02000806 were detected (Table IV). Most of the PAIs were previously identified from *Escherichia coli* [e.g., locus of enterocyte effacement (LEE)] and *Pseudomonas* (e.g., PAPI-1 and T-PAI).

larly interspaced short palindromic repeats.

Screening of antibiotic resistance genes. To reveal the genes relevant to the antibiotic resistance of MDR-SHH02, sequence alignment of the protein sequences of antibiotic resistance genes in ARDB and MDR-SHH02 genomic sequence was performed. Based on the selection criteria, a total of 12 gene

Table III. The top 10 predicted protein domains in the genomic sequence of *A. baumannii* MDR-SHH02.

Gene ID	Protein domain	P-value
MDRSHH02002231	AcrB/AcrD/AcrF family	<1.0E-300
MDRSHH02000985	Monomeric isocitrate dehydrogenase	<1.0E-300
MDRSHH02003056	AcrB/AcrD/AcrF family	<1.0E-300
MDRSHH02000840	AcrB/AcrD/AcrF family	<1.0E-300
MDRSHH02003594	AcrB/AcrD/AcrF family	<1.0E-300
MDRSHH02000667	AcrB/AcrD/AcrF family	<1.0E-300
MDRSHH02000153	Phosphoenolpyruvate carboxylase	2.70E-292
MDRSHH02000199	Urocanase	5.20E-287
MDRSHH02000831	Phosphoenolpyruvate carboxykinase	2.70E-279
MDRSHH02003134	AcrB/AcrD/AcrF family	5.50E-266

A. baumannii, Acinetobacter baumannii.

sequences (e.g., MDRSHH02002408, MDRSHH02000600 and MDRSHH02000597) in MDR-SHH02 were highly similar

Table IV. Pathogenicity islands homologous to a region in the genomic sequence of A. baumannii MDR-SHH02.

Gene ID	Start	End	Size (bp)	no. of ORFs	PAIs homologous to this region
MDRSHH02000806	332	21071	20740	18	PAPI-1 (Pseudomonas aeruginosa PA14)
					PAGI-2(C) (Pseudomonas aeruginosa C)
					PAGI-3(SG) (Pseudomonas aeruginosa SG17M)
					PAGI-5 (Pseudomonas aeruginosa PSE9)
					PPHGI-1 (Pseudomonas syringae pv. phaseolicola 1302A)
					SPI-7 (Salmonella enterica subsp. enterica serovar Typhi str. CT18)
					SPI-7 (Salmonella enterica subsp. enterica serovar Typhi Ty2)
					AbaR25 (Acinetobacter baumannii BJAB07104)
					AbaR26 (Acinetobacter baumannii BJAB0868)
					Hrp PAI (Erwinia amylovora 321)
					S-PAI (Pseudomonas cichorii 83-1)
					Hrp PAI (Pseudomonas syringae pv. tomato DC3000)
					Hrp PAI (Pseudomonas syringae pv. tomato str. DC3000)
					T-PAI (Pseudomonas viridiflava LP23.1a)
					T-PAI (Pseudomonas viridiflava PNA3.3a)
					S-PAI (Pseudomonas viridiflava RMX23.1a)
					S-PAI (Pseudomonas viridiflava ME3.1b)
					S-PAI (Pseudomonas viridiflava RMX3.1b)
					Not named (Corynebacterium urealyticum DSM 7109)
					LEE (Citrobacter rodentium DBS100)
					PAI I 536 (Escherichia coli 536)
					LEE (Escherichia coli E2348/69)
					LEE (Escherichia coli O157:H7 str. EDL933 ATCC43895)
					LEE (Escherichia coli 71074)
					LEE (Escherichia coli 83/39)
					LEE (Escherichia coli REPEC 84/110-1)
					LEE (Escherichia coli RW1374)
					LEE (Escherichia coli RDEC-1)
					LEE (Escherichia coli O157:H7 EDL933)
					LEE (Escherichia coli O157:H7 str. Sakai)
					LEE (Escherichia coli O103:H2 str. 12009)
				LEE (Escherichia coli O26:H11 str. 11368)	
				LEE (Escherichia coli O111:H- str. 11128)	
				LEE II (Escherichia coli 413/89-1)	
				AGI-3 (Escherichia coli BEN2908)	
					TAI (Escherichia coli O157:H7 EDL933)
				OI-122 (Escherichia coli O157:H7 EDL933)	
				PAI I CFT073 (Escherichia coli CFT073)	
				Not named (Escherichia coli UMN026)	
					SESS LEE (Salmonella enterica subsp. salamae serovar Sofia S1296
					SESS LEE (Salmonella enterica subsp. salamae serovar Sofia S1635
					SHI-1 (Shigella flexneri 2a str. 301)
					SHI-1 ( <i>Shigella flexneri</i> 2a str. 2457T)
					SRL (Shigella flexneri 2a YSH6000)
					Not named (Yersinia pestis CO92)

A. baumannii, Acinetobacter baumannii; ORF, open reading frame; PAIs, pathogenicity islands.

Table V. The potential gene sequences in A. baumannii MDR-SHH02 relevant to antibiotic resistance.

Gene ID	Resistance gene type from ARDB	Resistance gene class	Antibiotics  Astromicin, gentamicin, sisomicin	
MDRSHH02002408	aac(3)-Ia	Aminoglycoside N-acetyltransferase, which modifies aminoglycosides by acetylation		
MDRSHH02000600	aac(6')-Ib	Aminoglycoside N-acetyltransferase, which modifies aminoglycosides by acetylation	Amikacin, dibekacin, isepamicin, netilmicin, sisomicin, tobramycin	
MDRSHH02000597	ant(2")-Ia	Aminoglycoside O-nucleotidylyltransferase, which modifies aminoglycosides by adenylylation	Dibekacin, <b>gentamicin</b> , kanamycin, sisomicin, <b>tobramycin</b>	
MDRSHH02000611	ant(3")-Ia	Aminoglycoside O-nucleotidylyltransferase, which modifies aminoglycosides by adenylylation	Spectinomycin, streptomycin	
MDRSHH02001946	aph33ib	Aminoglycoside O-phosphotransferase, which modifies aminoglycosides by phosphorylation	Streptomycin	
MDRSHH02002406	aph(3')-Ia	Aminoglycoside O-phosphotransferase, which modifies aminoglycosides by phosphorylation	Gentamincin b, kanamycin, neomycin, paromomycin, lividomycin, ribostamycin	
MDRSHH02001945	aph(6)-Id	Aminoglycoside O-phosphotransferase, which modifies aminoglycosides by phosphorylation	streptomycin	
MDRSHH02000608	bl2b_tem, bl2b_tem1	Class A $\beta$ -lactamase. This enzyme breaks the $\beta$ -lactam antibiotic ring open and deactivates the molecule's antibacterial properties	Cephalosporin, penicillin, cephalosporin i, cephalosporin ii	
MDRSHH02000599	catb3	Group B chloramphenicol, acetyltransferase which can inactivate chloramphenicol. Also referred to as xenobiotic acetyltransferase	Chloramphenicol	
MDRSHH02000610	sul1	Sulfonamide-resistant dihydropteroate synthase, which can not be inhibited by sulfonamide	Sulfonamide	
MDRSHH02001738	sul2	Sulfonamide-resistant dihydropteroate synthase, which can not be inhibited by sulfonamide	Sulfonamide	
MDRSHH02001941	tetb	Major facilitator superfamily transporter, tetracycline efflux pump	Tetracycline	

A. baumannii, Acinetobacter baumannii; ARDB, Antibiotic Resistance Genes Database; DDA, disc diffusion assay. The antibiotics in bold are resisted by A. baumanii MDR-SHH02 in the DDA.

gentamicin, amikacin, tobramycin, spectinomycin, streptomycin and neomycin (Table V), which was partly consistent with the aforementioned results of antibiotic-resistance assay.

### Discussion

In the present study, we reported the draft genomic sequence of the clinical MDR *A. baumannii* strain, MDR-SHH02, and predicted one gene seuqence homologous to 45 PAIs and 12 potential gene sequences relevant to antibiotic resis-

tance. The antibiotic-resistance assay and the high similarity between the 12 gene sequences in MDR-SHH02 and the sequences of antibiotic resistance genes in ARDB, revealed that MDR-SHH02 was resistant to multiple antibiotics.

According to the prediction of PAIs, the gene sequence MDRSHH02000806 was homologous to 45 PAIs, such as LEE and PAPI-1. LEE PAIs were previously identified from multiple enteropathogenic *Escherichia coli* strains, and they are highly conserved in gene order and nucleotide sequence (25,26). PAPI-1 was previously identified from the *P. aeruginosa* 

strain, PA14, and it contributes directly and synergistically along with PAPI-2 to the virulence of PA14 (27). Therefore, the virulence of MDR-SHH02 may be due to the presence of MDRSHH02000806 homologous to these PAIs.

In this study, we discovered a set of gene sequences that were highly similar to the sequences of multiple genes encoding aminoglycoside-modifying enzymes (AMEs), including 2 aminoglycoside N-acetyltransferase genes [aac(3)-Ia and aac(6')-Ib], 2 aminoglycoside O-nucleotidylyltransferase genes [ant(2")-Ia and ant(3")-Ia], and 3 aminoglycoside O-phosphotransferase genes [aph33ib, aph(3')-Ia and aph(6)-Id]. The expression of AMEs enables bacteria to catalyze the modification of amino and hydroxyl groups on sugar moieties, such as aminoglycosides (28), which is a major cause of aminoglycoside resistance in many bacteria (29). The majority of aminoglycoside-resistant Acinetobacter isolates have the ability of enzymatic modification of aminoglycosides by acetyltransferases, nucleotidyltransferases and/or phosphotransferases (30). Previous studies have reported the prevalence of multiple AME genes [e.g., aac(3)-Ia, and aac(6')-Ib] in a set of A. baumannii isolates that are resistant to various aminogl ycosides (e.g., amikacin, gentamicin and tobramycin) (31,32). Besides, Aph(6)-Id and ant(3")-Ia have been detected in the A. baumannii strain, MRSN 12227, which is resistant to various antibiotics, such as amikacin, tobramycin and cefotaxime (33). Another study reported that ant(2")-Ia present in a group of A. baumannii isolates (62.6%) was associated with resistance to the tested aminoglycosides (amikacin, tobramycin and gentamicin) (34). Furthermore, in this study, we found that MDR-SHH02 was resistant to 19 antibiotics, such as several types of aminoglycosides (amikacin, gentamicin and tobramycin), indicating that the resistance of MDR-SHH02 to aminoglycosides likely resulted from the coding sequences highly similar to AME genes. However, aph33ib has not been previously detected in A. baumannii isolates, and it is worthy of further study. For example, following knockout and complementation of the gene sequence that is highly similar to aph33ib, the resistance of MDR-SHH02 to aminoglycosides is determined.

In this study, several gene sequences in MDR-SHH02 had a high similarity to class A β-lactamase genes (*bl2b\_tem* and *bl2b\_tem1*), group B chloramphenicol acetyltransferase gene (*catb3*), sulfonamide-resistant dihydropteroate synthase genes (*sul1* and *sul2*) and tetracycline efflux pump gene (*tetb*). The gene *bl2b\_tem* has been detected in *Staphylococcus aureus* (35), and *bl2b\_tem1* was detected in a series of marine bacteria, such as *Pelagibacter*, *Polaribacter* and *Roseobacter* (36). However, there is no evidence to support that *bl2b\_tem* and *bl2b\_tem1* are carried by *A. baumannii* isolates. All other genes (*catb3*, *sul1*, *sul2* and *tetb*) have been found in *A. baumannii* isolates (37-40).

Despite the aforementioned results, there were still several limitations to this study. The association of MDRSHH02000806 with the virulence of MDR-SHH02 needs to be validated in further studies. Besides, the associations between the 12 gene sequences similar to AME genes and the antibiotic resistance of MDR-SHH02 are also needs to be confirmed in further studies. We aim to conduct such experiments in our future studies.

In conclusion, this study fulfilled the draft genomic sequence of the clinical MDR *A. baumannii* strain, MDR-SHH02, and 12 gene sequences in MDR-SHH02 had a highly similarity

to the sequences of genes encoding AMEs [e.g., aac(3)-Ia, ant(2")-Ia, aph33ib and aph(3")-Ia], β-lactamase genes (bl2b\_tem and bl2b\_temI), sulfonamide-resistant dihydropteroate synthase genes (sul1 and sul2), catb3 and tetb. Of these genes, aph33ib, bl2b\_tem and bl2b\_temI were potential new antibiotic resistance genes. Furthermore, the antibiotic-resistance assay revealed that MDR-SHH02 was resistant to multiple antibiotics, such as amikacin, gentamicin and tobramycin. These findings were expected to enrich the data of antibiotic resistance genes in MDR A. baumannii, and provide a clinical guidance for the personalized therapy of A. baumannii-infected patients.

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