

# The increasing threat of silver-resistance in clinical isolates from wounds and burns

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**Purpose:** The widespread use of silver-containing compounds has led to emergence of silver-resistant bacteria. Few studies are available on the detectability of plasmid-mediated silver-resistance in developing countries. Therefore, we aimed to detect silver-resistance in isolates from wounds and burns, and to genetically characterize plasmid-mediated silver-resistance genes (*sil* genes).

**Methods:** One hundred and fifty clinical isolates were obtained from burns and wounds. They were identified using the suitable Analytical Profile Index and MicroScan identification systems. Their antimicrobial susceptibility was tested by the disk diffusion and broth microdilution methods. Their silver nitrate (AgNO<sub>3</sub>) minimum inhibitory concentration (MIC) was determined using the broth macrodilution method. The presence of different *sil* genes on plasmids extracted from silver-resistant isolates and the replicon types of the extracted plasmids were investigated using polymerase chain reaction (PCR). The ability of these plasmids to impart silver-resistance was tested by transformation.

**Results:** All except two isolates were multidrug-resistant. Nineteen silver-resistant bacterial isolates (12.6%) were detected; with AgNO<sub>3</sub> MIC ≥512 µg/mL. They were identified as *Klebsiella pneumoniae* (n=7), *Staphylococcus aureus* (n=4), *Escherichia coli* (n=2), *Enterobacter cloacae* (n=2), *Pseudomonas aeruginosa* (n=2) and *Acinetobacter baumannii* (n=2). PCR revealed the presence of different *sil* genes on the extracted plasmids. Plasmid transformation resulted in the transfer of silver-resistance to the resulting transformants. The extracted plasmids had different replicon types.

**Conclusion:** Plasmid-mediated silver-resistance was detected for the first time, in clinical *P. aeruginosa*, *A. baumannii* and *S. aureus* isolates; in addition to its detection in *K. pneumoniae*, *E. coli* and *Enterobacter cloacae*. Therefore, strict monitoring on the use of silver compounds in medical settings is required; with implementation of an approved standardized method for silver-resistance detection.

**Keywords:** multidrug-resistance, plasmid-mediated, replicon type, *sil* genes

## Introduction

The silver cation (Ag<sup>+</sup>) has been employed long ago as an antimicrobial agent (eg, AgNO<sub>3</sub> solution and silver sulfadiazine cream), for its microbicidal properties, minimum toxicity to human cells and therapeutic activity; especially in the prevention and treatment of burns and chronic wounds infections.<sup>1</sup>

Silver ions cause respiration inhibition, membrane damage and destruction of the proton motive force in sensitive organisms; through their interaction with thiol groups present in membrane proteins or enzymes and consequently the respiratory chain enzymes.<sup>2</sup> It can also generate reactive oxygen species that interfere with DNA replication<sup>3</sup> and increase membrane permeability.<sup>4</sup>

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Silver-impregnated dressings and antimicrobial coatings are used in infection management and stimulation of wound healing.<sup>5,6</sup> Also, silver is used as a component in dental amalgam and silver-impregnated medical devices, such as catheters and heart valves.<sup>7</sup> The widespread and uncontrolled use of silver may result in increasing the rate of bacterial resistance to silver-containing compounds where silver-resistant bacteria are as problematic as antibiotic-resistant ones.<sup>8</sup>

Acquired silver-resistance can be of endogenous (mutational) or exogenous (horizontally acquired) origin. Endogenous resistance to silver in Gram-negative bacteria can arise from derepression of the chromosomal *Cus* system and loss of outer membrane porins.<sup>9,10</sup> Also, derepression of the expression of silver-resistance genes (*sil* genes), integrated into *Escherichia coli* chromosome, has been reported to result in silver-resistance after stepwise adaptation in laboratory.<sup>11</sup> Staehlin and colleagues<sup>12</sup> have described a heavy metal homeostasis/resistance island, named as Copper homeostasis and silver resistance island (CHASRI), which was assembled and dispersed within members of *Enterobacteriaceae* family. Recently, CHASRI was found to be equally distributed in the plasmids and chromosomes of *Enterobacteriaceae* and was able to provide silver-resistance after mutation in *CusS* and/or *silS* genes of the CHASRI island.<sup>13</sup>

Exogenous silver-resistance was documented to be due to acquisition plasmid-encoded *sil* system.<sup>14,15</sup> The first encountered exogenous (horizontally acquired) silver-resistance was detected in a bacterial strain of *Salmonella enterica* serovar Typhimurium that caused an outbreak on a burns ward in 1975; causing the death of three patients.<sup>16</sup> Later, this resistance phenotype was detected in other bacterial species.<sup>17,18</sup> The first described plasmid was isolated from this *Salmonella* strain and named pMG101. The silver-resistance determinant comprises nine Open Reading Frames (ORFs) defining three transcriptional units (*silE*, *silS*, *silR*, *silC*, *silF*, *silB*, *silA*, ORF105 and *silP*).<sup>8</sup> These acquired silver-resistance genes are usually located on plasmids of the incompatibility group HI2 (IncHI2).<sup>19</sup>

Silver-resistance, whether endogenous or exogenous mediated, arises mainly from restriction of intracellular accumulation of silver. *silE* is a small periplasmic silver-binding protein that binds silver specifically at the cell surface; presenting the first line of resistance against silver toxicity. Also, there is a two-component silver-responsive transcriptional regulation system; *silRS*. This system controls the expression of a silver efflux ATPase; *silP*, the tripartite

*silCBA* silver effluxer and *silF*, which is periplasmic silver chaperone.<sup>8,20</sup> Endogenous silver-resistance, due to porin loss and over expression of *Cus* system that causes silver efflux, also limits the accumulation of silver inside the cell.<sup>14</sup>

Detection of exogenously acquired silver-resistance in *K. pneumoniae* and *Enterobacter* spp. isolates is highly worrisome where only few therapeutic options will be available for treatment of these bugs; if any. The available data on exogenously acquired silver-resistance and the distribution of *sil* genes among hospital pathogens, in developing countries; including Egypt, are rare. Therefore, the aim of this study was to detect silver-resistance and the associated resistance to other antibiotics as well as to detect and characterize plasmid-encoded *sil* genes; in different clinical isolates from wounds and burns infections.

## Materials and methods

### Analysis of silver-resistance genes in different bacterial species

*sil* genes present in the publically available genomes, of different bacterial species, were retrieved from Patricbrc web resources (<https://www.patricbrc.org>).<sup>21</sup> The status of retrieved *sil* genes, whether chromosomal or plasmid encoded, was confirmed by searching their accession numbers on NCBI nucleotide database.

### Microorganisms

One hundred and fifty bacterial isolates from wounds (n=102) and burns (n=48) were collected between October 2015 and April 2016 from the Central Laboratories, El-Demerdash Hospital, Ain Shams University, Cairo, Egypt. All wound and burn sepsis samples, collected during this period, were sent to laboratories for routine diagnosis. They were collected using sterile cotton swabs and transferred to the laboratories in Cary-Blair transport system. In the laboratory, they were inoculated aseptically onto blood and MacConkey agar plates (Oxoid Ltd., Hampshire, UK) and incubated at 35–37°C for 24–48 hrs. The isolated colonies were identified using the suitable Analytical Profile Index (Biomérieux Inc., France) and MicroScan (Beckman Coulter, USA) identification systems; following manufacturers' instructions. *Escherichia coli* TOP10 was used as a recipient strain in the transformation experiments.

### Ethics approval and informed consent

This study was approved by the ethical committee of the Faculty of Pharmacy, Cairo University, Cairo, Egypt

(approval number: MI 1537). An oral informed consent was obtained from the patients since the samples were collected during routine diagnosis and no special interventions were made for the research purpose.

## Antimicrobial susceptibility testing of the collected isolates

The antimicrobial susceptibility testing of the isolated bacterial species was determined by Kirby-Bauer disk diffusion method<sup>22</sup> except against vancomycin and polymyxin B where antimicrobial susceptibility was determined by the broth microdilution method.<sup>23</sup> The selection of the antimicrobial agents was based on the type of the organism being tested, and the results were interpreted following the Clinical and Laboratory Standards Institute (CLSI) and European Committee on Antimicrobial Susceptibility Testing (EUCAST) interpretive criteria.<sup>24,25</sup> The antibiotic disks included in the study were gentamicin 10 µg, piperacillin/tazobactam 100 µg/10 µg, imipenem 10 µg, meropenem 10 µg, cefazolin 30 µg, ceftazidime 30 µg, cefepime 30 µg, ceftiofur 30 µg, ciprofloxacin 5 µg, sulfamethoxazole/trimethoprim 23.75 µg/1.25 µg, tigecycline 15 µg, aztreonam 30 µg, chloramphenicol 30 µg, doxycycline 30 µg, fusidic acid 10 µg, clindamycin 2 µg, erythromycin 15 µg, linezolid 30 µg and quinupristin/dalfopristin 15 µg (All from Oxoid Ltd., Hampshire, UK).

## Determination of AgNO<sub>3</sub> minimum inhibitory concentration (MIC)

The susceptibility of different isolates to AgNO<sub>3</sub> (Sigma-Aldrich, USA) was evaluated by MIC determination using broth macrodilution method.<sup>23</sup> Two-fold serial dilutions of AgNO<sub>3</sub> were prepared, using deionized water, to give a concentration range from 4 to 512 µg/mL. Isolates with MIC of ≥512 µg/mL, against AgNO<sub>3</sub>, were classified as silver-resistant.<sup>26</sup>

## Detection of different *sil* genes

Genomic DNA was extracted from all isolates by the boiling method.<sup>27</sup> The extracted DNA was tested for the presence of *silP*, *silB*, *silF*, *silE*, *silA*, *silCBA* and *silRS* genes by the polymerase chain reaction (PCR) as suggested by Woods et al,<sup>17</sup> with minor modifications in primers' sequences to increase the specificity of the reaction (Table 1). Also, reaction temperatures were optimized using gradient PCR. The identity of the positive amplicons, from each reaction, was confirmed by sequencing and isolates that were confirmed to possess the tested genes were used as positive controls for subsequent reactions.

PCR products were visualized on 1.5% agarose gel (Axygen Biosciences, CA, USA) by electrophoresis using 100-bp ladder (Invitrogen, ThermoFisher Scientific, MA, USA). Randomly selected PCR products, of the expected size, were purified with QIAquick PCR Purification Kit (Qiagen, Crawley, UK) and sequenced by ABI<sup>TM</sup>3500 Genetic Analyzer DNA sequencer (Applied Biosystems, Foster City, CA). Similarity searches for the nucleotide sequences were performed with BLASTN program (<http://www.ncbi.nlm.nih.gov/blast>) using default settings.

## Determination of the origin of *sil* genes

Plasmids were extracted from all isolates, that were positive for any of the tested *sil* genes, using a GeneJET Plasmid Miniprep kit (Invitrogen, ThermoFisher Scientific, MA, USA); following the manufacturer's instructions. The extracted plasmids were tested for the presence of different *sil* genes; as described previously.

## Confirmation of the ability of extracted plasmids to impart silver-resistance phenotype

The ability of the extracted plasmids to confer silver-resistance was tested by transformation into *E. coli* TOP10.

**Table 1** Primers used for *sil* genes detection with the corresponding annealing temperatures and the expected amplicon size

Gene	Forward primer	Reverse primer	Annealing temperature (°C)	Amplicon size (bp)
<i>silE</i>	aggggaaacggtctgacttc	atatccatgagcgggtcaac	52	221
<i>silB</i>	caaagaacagcgcgtgatta	gctcagacattgctgcata	52	233
<i>silP</i>	cctgggtttacagcgtcatt	atggcacctgaggtttgttc	52	175
<i>silF</i>	cgatatgaatgctgccagt	attgccctgctgaataaacg	52	229
<i>silA</i>	cttgagcatccaacaagaa	ctgccagtacaggaacct	54	162
<i>silRS</i>	gacggcaatcgcaatcagatt	gtggaggatactgcgagagc	54	192
<i>silCBA</i>	cgggaaacgctgaaaaattatgaa	gtacgttcccagcaccagtt	54	189

Briefly, the extracted plasmids from each silver-resistant bacterial isolate were electroporated into electrocompetent *E. coli* TOP10 cells using a Gene Pulser electroporator (Gene MicroPulser™ Electroporation System, Bio-Rad, USA); according to the manufacturer's instructions. Transformants were selected on LB agar (Merk Millipore, Germany) plates supplemented with 64 µg/mL AgNO<sub>3</sub> (The AgNO<sub>3</sub> MIC of *E. coli* TOP10 was 32 µg/mL). Transformants were tested for their antimicrobial susceptibility, to different antibiotics, using Kirby-Bauer disk diffusion method and broth microdilution method; as mentioned before. Their AgNO<sub>3</sub> MICs were determined by broth microdilution method, as described previously.

## PCR-based plasmid replicon typing

Plasmid replicon typing, of the plasmids extracted from the silver-resistant isolates, was performed according to Johnson et al.<sup>28</sup>

## Results

### Bioinformatics analysis of silver-resistance genes

*sil* genes were detected in 400 different bacterial genomes present in the publically available genomes' sector of Patric resources. The plasmid or chromosomal location, of the retrieved genomes containing *sil* genes, was determined. Twenty one genomes were confirmed to have plasmid-encoded *sil* genes. These genomes were from *Klebsiella pneumoniae* (seven strains), *Escherichia coli* (nine strains) and *Enterobacter cloacae* (five strains). All plasmid-encoded *sil* cassettes contained the nine open reading frames described by Gupta et al.<sup>8</sup> No *sil* genes were detectable in the publically available genomes of *Acinetobacter baumannii* or *Staphylococcus aureus*.

### Identification of bacterial isolates from burns and wounds

Bacterial isolates collected from wounds and burns were identified as *K. pneumoniae* (n=32), *E. coli* (n=22), *P. aeruginosa* (n=21), *Enterobacter cloacae* (n= six isolates), *Klebsiella aerogenes* (n= two isolates), *A. baumannii* (n=14), *Proteus mirabilis* (n=15) and *S. aureus* (n=38).

### Antimicrobial and silver susceptibility patterns of the collected isolates

The antibiogram of the isolated bacterial species revealed that all except two isolates were multidrug-resistant

“MDR” according to the definition of Magiorakos et al.<sup>29</sup> MDR is defined as non-susceptibility to at least one agent in three or more antimicrobial categories.<sup>29</sup> *E. coli* TOP10 was susceptible to all tested antimicrobials. The results of identification of isolated bacterial species and their antibiogram are given in Table S1.

Of the 150 tested isolates, nineteen bacterial isolates were silver-resistant; with a MIC ≥512 µg/mL; against AgNO<sub>3</sub>. They were identified as *K. pneumoniae* (seven isolates), *E. coli* (two isolates), *Enterobacter cloacae* (two isolates), *P. aeruginosa* (two isolates), *A. baumannii* (two isolates) and *S. aureus* (four isolates). None of the tested *Klebsiella aerogenes* or *Proteus mirabilis* was resistant to AgNO<sub>3</sub>. The silver-susceptible bacterial isolates had MIC range of 32–128 µg/mL; against AgNO<sub>3</sub>.

### Detection of the silver-resistance genes

All isolates were tested for the presence of different *sil* genes using seven primer pairs capable of detecting the eight identified silver-resistance genes. Silver-susceptible isolates were negative for all tested *sil* genes; however, all silver-resistant isolates recorded positive results for at least three of the tested *sil* genes. All tested *sil* genes were detectable in six out of the 19 silver-resistant isolates (three *K. pneumoniae* isolates, one *E. coli* isolate, one *A. baumannii* isolate and one *S. aureus* isolate). Other isolates were positive only for some but not all of the tested *sil* genes. Table 2 describes the detectability of different *sil* genes in each of the tested isolates. The obtained sequences, of the selected PCR amplicons, were deposited in GenBank under accession numbers MH229478 to MH229484. *E. coli* TOP10 was negative for all tested *sil* genes.

### Origin of silver-resistance genes and plasmid replicon typing

All detected *sil* genes were found to be encoded on the extracted plasmids. To further confirm the location of the detected *sil* genes on the extracted plasmids and not on chromosomal contaminant, the DNA was extracted from the plasmid bands separated on the agarose gel and tested for the presence of different *sil* genes. The results confirmed the plasmid location of *sil* genes. Transformation of the plasmids extracted from silver-resistant isolates into *E. coli* TOP10 resulted in 19 bacterial transformants with the same silver-resistance level as that of the original isolates (AgNO<sub>3</sub> MIC ≥512 µg/mL). The results of the antimicrobial susceptibility testing of the transformants revealed susceptibility to all of the

**Table 2** Detectability of *sil* genes in silver-resistant isolates and the obtained transformants

Bacterial species Tested genes	<i>silRS</i>	<i>silE</i>	<i>silCBA</i>	<i>silB</i>	<i>silA</i>	<i>silP</i>	<i>silF</i>
176K	+	-	+	+	+	+	+
176K <sub>T</sub>	+	-	+	+	+	+	+
197K	+	-	+	+	+	+	+
197K <sub>T</sub>	+	-	+	+	+	+	+
199K	+	+	+	+	+	+	+
199K <sub>T</sub>	+	+	+	+	+	+	+
215K	+	+	+	+	+	+	+
215K <sub>T</sub>	+	+	+	+	+	+	+
465K	+	-	+	+	+	+	+
465K <sub>T</sub>	+	-	+	+	+	+	+
724K	+	-	+	+	+	+	+
724K <sub>T</sub>	+	-	+	+	+	+	+
793K	+	+	+	+	+	+	+
793K <sub>T</sub>	+	+	+	+	+	+	+
205E	+	+	+	+	+	+	+
205E <sub>T</sub>	+	+	+	+	+	+	+
263E	+	+	-	+	+	+	-
263E <sub>T</sub>	+	-	-	+	+	+	-
285En	+	-	-	+	+	-	-
285En <sub>T</sub>	+	-	-	+	+	-	-
367En	+	-	-	+	+	-	-
367En <sub>T</sub>	+	-	-	+	+	-	-
314P	+	-	-	-	+	+	-
314P <sub>T</sub>	+	-	-	-	+	-	-
646P	+	+	-	-	+	+	-
646P <sub>T</sub>	+	+	-	-	+	-	-
407Ac	+	+	+	+	+	+	+
407Ac <sub>T</sub>	+	+	+	+	+	+	+
457Ac	-	+	-	+	+	+	+
457Ac <sub>T</sub>	-	+	-	+	+	+	+
447S	+	+	+	+	+	+	+
447S <sub>T</sub>	+	+	+	+	+	+	+
579S	-	-	+	+	+	+	+
579S <sub>T</sub>	-	-	+	+	+	+	+
586S	-	-	-	+	-	+	+
586S <sub>T</sub>	-	-	-	+	-	+	+
601S	-	+	-	-	+	+	-
601S <sub>T</sub>	-	+	-	-	+	+	-

**Notes:** Dark shaded cells indicate the genes that were not detectable in the some of the resulting transformants.

**Abbreviation:** T, the corresponding transformant.

tested antimicrobial agents. They exhibited resistance pattern similar to that of the *E. coli* TOP10, against these agents. The resulting transformants possessed the same *sil* genes as that in

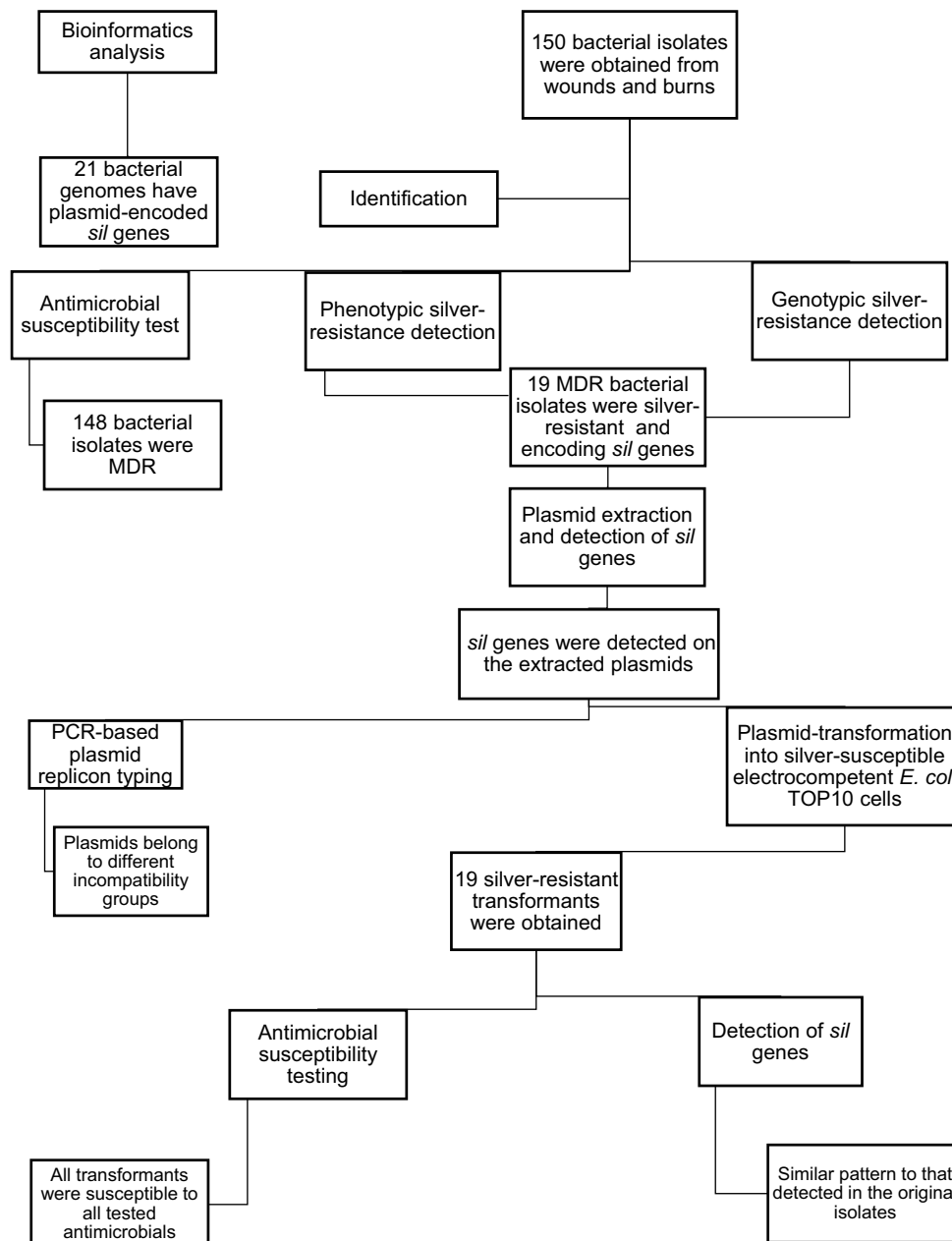
the original isolates, except three transformants where transformant 263E<sub>T</sub> was negative for *silE* and transformants 314P<sub>T</sub> and 646P<sub>T</sub> were negative for *silP* (Table 2).

Plasmid replicon typing was performed on the extracted plasmids. Most of the plasmids belonged to the incompatibility groups IncX (four isolates; 215K, 465K, 447S and 586S), IncHI1 (three isolates; 199K, 724K and 457Ac), IncN (three isolates; 205E, 285En and 367En), IncI1 (three isolates; 798K, 314P and 646P) and IncB/O (two isolates; 579S and 601S). In addition, one plasmid belonged to each of the incompatibility groups IncFrep (176K), IncFIC (197K), IncP (263E) and IncA/C (407Ac).

The plan of work and the obtained results are summarized in Figure 1.

## Discussion

Silver has been considered as an alternative to commonly used antimicrobials due to the widespread antimicrobial resistance.<sup>30</sup> However, appearance of plasmid-mediated silver-resistance presented an alarm where silver-resistance can limit the effectiveness of silver-based



**Figure 1** Scheme of the work process with the obtained results.

**Abbreviations:** MDR, multidrug-resistant; *sil* genes, silver-resistance genes.

antimicrobials in the future. Only few studies addressed this threat; especially in developing countries.

In this study, 150 clinical isolates, from wounds and burns, were collected. Most of the isolated species were Gram-negative bacteria (75%), which comes in agreement with previous studies about the pattern of bacterial types colonizing wounds and burns.<sup>31,32</sup> *S. aureus* was the predominantly detected organism (26%) followed by *K. pneumoniae* (21%). Some previous studies have reported *S. aureus* as the predominant isolate from wounds.<sup>33,34</sup> On the other hand, Srinivasan et al<sup>35</sup> and Forson et al<sup>36</sup> detected *K. pneumoniae* and *P. aeruginosa* as the predominant organisms, respectively. This difference, in predominance, may have resulted from variation in the applied treatment practices.

It was difficult to compare our silver-resistance rate to that published in most of available studies; where different cutoff values were used for defining silver-resistant isolates.<sup>14,37</sup> Since no universal standard MIC breakpoint, for silver-resistance, is available, we hereby take this opportunity to highlight the urgent need for defining the standard MIC breakpoint for silver-resistance. Only two studies were concerned with silver-resistance rates, in clinical isolates, and used the same MIC breakpoint as that we used here. We were able to detect silver-resistance in 12.6% of the collected isolates. Similarly, Sütterlin and coworkers<sup>38</sup> detected phenotypic silver-resistance in six out of 56 bacterial isolates (11%). Also, in another study, Sütterlin and coworkers<sup>26</sup> recorded phenotypic silver-resistance in 13% of *Enterobacter* spp., and 3% of *Klebsiella* spp. out of 752 blood stream isolates.

What worsens the condition is that all of our isolated silver-resistant bacteria were MDR. This is not surprising where most of our silver-resistant isolates belonged to ESKAPE pathogens (*Enterococcus faecium*, *Staphylococcus aureus*, *Klebsiella pneumoniae*, *Acinetobacter baumannii*, *Pseudomonas aeruginosa* and *Enterobacter* spp.) which are known with their MDR phenotype.<sup>39</sup> They were also detectable in *E. coli* where MDR phenotypes, in different clones of *E. coli*, isolated from extraintestinal infection, has been reported previously.<sup>40</sup> However, it is highly worrisome regarding the limited therapeutic options available; if any.<sup>41</sup>

Silver-resistance was detected in *K. pneumoniae* (seven isolates), *E. coli* (two isolates), *P. aeruginosa* (two isolates), *A. baumannii* (two isolates), *Enterobacter cloacae* (two isolates) and *S. aureus* (four isolates). The detection of silver-resistance among different Gram-negative bacteria belonging to *Enterobacteriaceae* family is common.<sup>38</sup> However, we are

reporting for the first time the detection of silver-resistant *P. aeruginosa*, *A. baumannii* and *S. aureus* clinical isolates. To our knowledge, only two studies were concerned with silver-resistance in *S. aureus* where Sütterlin and colleagues<sup>38</sup> recorded their AgNO<sub>3</sub> MIC to be in the range of 16–32 µg/mL (sensitive range according to the used MIC breakpoint). Also, in a study carried by Randall and coworkers,<sup>42</sup> no evidence was found for silver-resistance in 876 clinical staphylococcal isolates, nor could any reduction in silver-susceptibility be selected upon extended exposure (42 days) of *S. aureus* to silver *in vitro*.

The cause for this variation, in the detectability of silver-resistance, is not clear. Our isolates were collected from wound and burn patients that were treated with silver sulfadiazine (1%) cream during the period of samples collection. The same was reported by Randall et al<sup>42</sup> about the use of silver compounds during the period of collection of their burn and diabetic foot ulcer *S. aureus* isolates (They were at least 230 isolates). Silver-resistance in our *S. aureus* isolates may have been resulted from the regular use of silver sulfadiazine cream in Egyptian facilities which may have selected for resistant isolates where different protocols may be applied in other areas; limiting the spread of resistance.

We searched the publically available genomes, of Patric resources, for plasmid-mediated *sil* genes; to check the possibility of their presence as part of sequenced plasmids. This was done to avoid any possibility of missing *sil* genes that may be present in species other than those available in literature. The detectability of plasmid-mediated *sil* genes, among our collection of both Gram-negative and positive isolates, was different from that available in Patric resources. In publically available genomes, plasmid-mediated *sil* genes were detectable in *K. pneumoniae*, *E. coli* and *Enterobacter cloacae* only. However, here we were able to detect these genes also in *A. baumannii*, *P. aeruginosa* and *S. aureus*. Previous studies reported the detection of plasmid-mediated *sil* genes only in clinical isolates from *Enterobacteriaceae* family as *Salmonella enterica* serovar Typhimurium isolated from burns ward,<sup>8</sup> *E. coli*,<sup>19</sup> *Enterobacter cloacae* and *K. pneumoniae*.<sup>14</sup>

Presence of plasmid-mediated *sil* genes, in isolates from wounds and burns, is highly dangerous. This is because of the possibility of plasmid transfer between different bacterial species colonizing the wound; where chronic wounds are usually colonized with a collection of different bacteria. They, therefore, provide an ultimate environment for plasmids to travel among contaminating

strains; especially that biofilms are becoming significant in this area.<sup>43</sup>

In our study, and in accordance with the detectability of different *sil* genes on plasmids from the publically available genomes in Patric resources, six isolates were positive for all tested *sil* genes. However, the remaining isolates missed one or more genes. Similar findings, regarding the absence of some *sil* genes, were reported previously.<sup>14,38,44</sup> In our study, isolates belonging to *Enterobacteriaceae* family mostly lacked *silE*, *silCBA* and *silF* genes. This is different from the gene possession described by Gupta et al<sup>19</sup> where only *silE* and *silP* genes were sometimes missed in *Enterobacteriaceae* members. No records are available about the *sil* genes possession in other bacterial species.

This difference in *sil* genes detectability may have resulted from the molecular method employed in gene detection; where all retrieved genes from the database were from a fully sequenced plasmid. In our study, as well as in others, the genes were detected mainly by PCR; where variation in gene sequences may have caused the failure of amplification by the used primers. Minor sequence variation ( $\approx 4\%$ ) was previously recorded in different *sil* genes.<sup>10,19</sup>

The possession of *sil* genes is not enough to account for phenotypic resistance to silver as derepression of Sil system is required for expression of silver-resistance phenotype.<sup>10</sup> Therefore, the ability of the extracted plasmids to impart silver-resistance was confirmed by transformation; where all the resulting transformants had the same level of silver-resistance. In addition, they have the same *sil* genes profile except three transformants. The reason underlying this variation in gene profiles is unclear.

Several cellular mechanisms can also account for silver-resistance, in addition to the acquisition of derepressed plasmid-encoded *sil* genes; as mutations in chromosomal Cus system together with the loss of outer membrane porins.<sup>9,10</sup> However, it is unlikely to be responsible for silver-resistance in all our transformants; as mutations causing silver-resistance were not detectable, in laboratory testing, even after several passages in suprainhibitory concentrations of silver nitrate.<sup>10</sup>

Plasmid replicon typing, of the extracted plasmids, was performed using the method of Johnson et al<sup>28</sup> for detection of incompatibility groups of plasmids from *Enterobacteriaceae* family. We applied this method to plasmids extracted from different bacterial species, whether *Enterobacteriaceae* or not, based on the history of the detectability of plasmid-mediated silver-resistance,

in clinical isolates from members of *Enterobacteriaceae* family only, as discussed previously.

All extracted plasmids were typable and carried only one of the tested replicon types; even those from organisms not belonging to *Enterobacteriaceae* family. This indicated that all of these plasmids may have originated from organisms belonging to the family *Enterobacteriaceae*. The transfer of plasmids from *Enterobacteriaceae* members to other Gram-negative or Gram-positive bacteria; such as *P. aeruginosa*, *A. baumannii* and *S. aureus*, has been reported previously.<sup>45-47</sup>

Silver-resistance genes are known to be carried only on plasmids of IncHI group.<sup>15,19</sup> However, our plasmid-mediated *sil* genes were detected on plasmids of different replicon types, as IncFrep, IncFIC, IncHI1, IncX, IncI1, IncN, IncP, IncA/C and IncB/O. This may have resulted from the integration of *sil* cassette into different plasmid types, mediated by Tn-7 transposition, as described by Randall et al.<sup>10</sup>

It was surprising that none of the obtained transformants gained resistance to any of the tested antimicrobials; to which the original isolates were resistant. This indicated that the antimicrobial resistance determinants, in our collected isolates, were not encoded on the same plasmid together with the detected silver-resistance genes which is quite opposite to the MDR phenotype associated with the plasmid pMG101.<sup>48</sup>

Further study is still required to investigate different mechanisms underlying this MDR phenotype and its possible linkage to silver-resistance. The results of plasmid replicon typing confirmed the presence of only one replicon type in each extracted plasmid preparation. However, only replicons from *Enterobacteriaceae* family were tested as indicated previously; where plasmids from other species may be present and go undetectable by *Enterobacteriaceae* plasmid replicon typing. Antimicrobial resistance determinants may have been encoded on large plasmids that are not transformed efficiently by electroporation.<sup>49</sup> Also, the mechanisms underlying MDR phenotype may not be plasmid encoded; as in case of mutations causing efflux pumps overexpression or loss of outer membrane porins or may be encoded by chromosomal genes.

This study is alarming regarding the spread of phenotypic silver-resistance and *sil* genes in species where they were not detectable before. However, several questions are still required to be answered, as few numbers of isolates were studied in each species. So, other studies are required to address the detectability of silver-resistance in large number of isolates from each species together with their



genetic background and clonal dissemination. In addition, nearly all of our isolated strains were MDR so it was difficult to draw a solid conclusion about the association between silver-resistance and MDR.

## Conclusion

We are reporting here, for the first time, the detectability of silver-resistance and plasmid-mediated *sil* genes in different types of clinical isolates (*P. aeruginosa*, *A. baumannii* and *S. aureus*). This is highly worrisome in terms of resistance spread; especially within the poly-microbial-infected wounds and burns, and the limited therapeutic options where all silver-resistant isolates were MDR. Therefore, a careful monitoring on the use of silver-containing antimicrobial agents is highly required with the development of an approved standardized method for the detection of silver-resistance.

## Disclosure

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest in regard to this work.

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# Supplementary material

**Table S1** Antimicrobial susceptibility pattern and the drug-resistance phenotype of different bacterial isolates

Isolate number	Identification	CN	TZP	IMP	MEM	KZ	CAZ	FEP	FOX	CIP	SXT	TGC	ATM	C	PB <sup>a</sup>	DO	FD	V <sup>a</sup>	DA	E	LZD	QD	DR			
176	<i>Escherichia coli</i> TOP10																							MDR		
177																									MDR	
179																										MDR
197																										MDR
198																										MDR
199																										MDR
204																										MDR
205																										MDR
215																										MDR
225																										MDR
227																										MDR
241																										MDR
270																										MDR
290																										MDR
298																										MDR
370																										MDR
371																										MDR
407																										MDR
416																										MDR
464																										MDR
465																										MDR
476																										MDR
492																										MDR
580																										MDR
646																										MDR
648																										MDR
724																										MDR
729																									MDR	
749																									MDR	
769																									MDR	
793																									MDR	
808																									MDR	

*Klebsiella pneumoniae* (K)

(Continued)

Table S1 (Continued).

Isolate number	Identification	CN	TZP	IMP	MEM	KZ	CAZ	FEP	FOX	CIP	SXT	TGC	ATM	C	PB <sup>a</sup>	DO	FD	V <sup>b</sup>	DA	E	LZD	QD	DR	
53	<i>Escherichia coli</i> (E)																						MDR	
197																								MDR
205																								MDR
211																								MDR
215																								MDR
229																								MDR
244																								MDR
263																								MDR
269																								MDR
276																								MDR
289																								MDR
290																								MDR
298																								MDR
315																								MDR
416																								MDR
437																								MDR
445																								MDR
466																								MDR
580																								MDR
599																								MDR
647																								MDR
682																								MDR
179		<i>Klebsiella aerogenes</i>																						MDR
314																								MDR
228																								MDR
285		<i>Enterobacter cloacae</i> (En)																						MDR
367																								MDR
465																								MDR
637																							MDR	
648																							MDR	

(Continued)

**Table S1** (Continued).

Isolate number	Identification	CN	TZP	IMP	MEM	KZ	CAZ	FEP	FOX	CIP	SXT	TGC	ATM	C	BP*	DO	FD	V*	DA	E	LZD	QD	DR	
24	<i>Proteus mirabilis</i>	■																					MDR	
176			■							■														MDR
199																								MDR
227																								MDR
228																								-
269																								MDR
288																								MDR
371																								MDR
463																								MDR
465																								MDR
477																								MDR
745																								MDR
749																								MDR
793																								MDR
817																								MDR

(Continued)

**Table S1 (Continued).**

Isolate number	Identification	CN	TZP	IMP	MEM	KZ	CAZ	FEP	FOX	CIP	SXT	TGC	ATM	C	PP*	DO	FD	V*	DA	E	LZD	QD	DR	
188	<i>Pseudomonas aeruginosa</i> (P)	■	■			■	■	■	■	■	■	■		■	■	■	■	■	■	■	■	■	MDR	
198		■	■			■	■	■	■	■	■	■		■	■	■	■	■	■	■	■	■	MDR	
199		■	■			■	■	■	■	■	■	■		■	■	■	■	■	■	■	■	■	■	MDR
227		■	■			■	■	■	■	■	■	■		■	■	■	■	■	■	■	■	■	■	MDR
232		■	■			■	■	■	■	■	■	■		■	■	■	■	■	■	■	■	■	■	MDR
236		■	■			■	■	■	■	■	■	■		■	■	■	■	■	■	■	■	■	■	MDR
241		■	■			■	■	■	■	■	■	■		■	■	■	■	■	■	■	■	■	■	MDR
269		■	■			■	■	■	■	■	■	■		■	■	■	■	■	■	■	■	■	■	MDR
289		■	■			■	■	■	■	■	■	■		■	■	■	■	■	■	■	■	■	■	MDR
314		■	■			■	■	■	■	■	■	■		■	■	■	■	■	■	■	■	■	■	MDR
365		■	■			■	■	■	■	■	■	■		■	■	■	■	■	■	■	■	■	■	MDR
400		■	■			■	■	■	■	■	■	■		■	■	■	■	■	■	■	■	■	■	MDR
417		■	■			■	■	■	■	■	■	■		■	■	■	■	■	■	■	■	■	■	MDR
438		■	■			■	■	■	■	■	■	■		■	■	■	■	■	■	■	■	■	■	MDR
441		■	■			■	■	■	■	■	■	■		■	■	■	■	■	■	■	■	■	■	MDR
465		■	■			■	■	■	■	■	■	■		■	■	■	■	■	■	■	■	■	■	MDR
492		■	■			■	■	■	■	■	■	■		■	■	■	■	■	■	■	■	■	■	MDR
599		■	■			■	■	■	■	■	■	■		■	■	■	■	■	■	■	■	■	■	MDR
646		■	■			■	■	■	■	■	■	■		■	■	■	■	■	■	■	■	■	■	MDR
749		■	■			■	■	■	■	■	■	■		■	■	■	■	■	■	■	■	■	■	MDR
808	■	■			■	■	■	■	■	■	■		■	■	■	■	■	■	■	■	■	■	MDR	

(Continued)

**Table S1** (Continued).

Isolate number	Identification	CN	TZP	IMP	MEM	KZ	CAZ	FEP	FOX	CIP	SXT	TGC	ATM	C	PB <sup>a</sup>	DO	FD	V <sup>b</sup>	DA	E	LZD	QD	DR	
1	<i>Acinetobacter baumannii</i> (Ac)																						MDR	
198																								MDR
203																								MDR
289																								MDR
407																								MDR
417																								MDR
418																								MDR
419																								MDR
457																								MDR
576																								MDR
592																								MDR
637																								MDR
818																								MDR
822																								MDR

(Continued)

Table S1 (Continued).

Isolate number	Identification	CN	TZP	IMP	MEM	KZ	CAZ	FEP	FOX	CIP	SXT	TGC	ATM	C	PB <sup>a</sup>	DO	FD	V <sup>a</sup>	DA	E	LZD	QD	DR	
48	<i>Staphylococcus aureus</i>																						MDR	
212																								MDR
244																								MDR
262																								MDR
264																								MDR
269																								MDR
278																								MDR
280																								MDR
281																								MDR
284																								MDR
285																								MDR
294																								MDR
304																								MDR
316																								MDR
338																								MDR
350																								MDR
365																								MDR
366																								MDR
399																								MDR
407																								MDR
410																								MDR
438																								MDR
443																								MDR
446																								MDR
447																								MDR
550																								MDR
578																							MDR	
579																							MDR	
581																							MDR	
584																							MDR	
586																							MDR	
599																							MDR	
601																							MDR	
607																							MDR	
611																							MDR	
612																							MDR	
648																							MDR	
661																							MDR	

**Notes:** Black cells, resistance to tested antibiotic; white cells, sensitive to tested antibiotic; light gray cells, intermediate result to tested antibiotic; Diagonal shaded cells, non-applicable antibiotics. <sup>a</sup>Susceptibility to vancomycin and polymyxin B was determined by MIC.

**Abbreviations:** CN, gentamicin; TZP, piperacillin/tazobactam; IMP, imipenem; MEM, meropenem; KZ, ceftazidime; CAZ, ceftazidime; FEP, cefepime; FOX, ceftioxiacin; SXT, sulfamethoxazole/trimethoprim; TGC, tigecycline; ATM, aztreonam; C, chloramphenicol; PB, polymyxin B; DO, doxycycline; FD, fusidic acid; V, vancomycin; DA, clindamycin; E, erythromycin; LZD, linezolid; QD, quinupristin/dalfopristin; DR, drug-resistance phenotype; MDR, Multidrug resistance.



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