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ORIGINAL RESEARCH

Consequences Of Long-Term Bacteria's Exposure To Silver Nanoformulations With Different PhysicoChemical Properties

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Purpose: Resistance to antibiotics is a major problem of public health. One of the alternative therapies is silver – more and more popular because of nanotechnology development and new possibilities of usage. As a component of colloid, powder, cream, bandages, etc., nanosilver is often recommended to treat the multidrug-resistant pathogens and we can observe its overuse also outside of the clinic where different physicochemical forms of silver nanoformulations (e.g. size, shape, compounds, surface area) are introduced. In this research, we described the consequences of long-term bacteria exposure to silver nanoformulations with different physicochemical properties, including changes in genome and changes of bacterial sensitivity to silver nanoformulations and/or antibiotics. Moreover, the prevalence of exogenous resistance to silver among multidrug-resistant bacteria was determined.

Materials and Methods: Gram-negative and Gram-positive bacteria strains are described as sensitive and multidrug-resistant strains. The sensitivity of the tested bacterial strains to antibiotics was carried out with disc diffusion methods. The sensitivity of bacteria to silver nanoformulations and development of bacterial resistance to silver nanoformulations has been verified via determination of the minimal inhibitory concentrations. The presence of *sil* genes was verified via PCR reaction and DNA electrophoresis. The genomic and phenotypic changes have been verified via genome sequencing and bioinformatics analysis.

Results: Bacteria after long-term exposure to silver nanoformulations may change their sensitivity to silver forms and/or antibiotics, depending on the physicochemical properties of silver nanoformulations, resulting from phenotypic or genetic changes in the bacterial cell. Finally, adaptants and mutants may become more sensitive or resistant to some antibiotics than wild types.

Conclusion: Application of silver nanoformulations in the case of multiple resistance or multidrug-resistant bacterial infection can enhance or decrease their resistance to antibiotics. The usage of nanosilver in a clinic and outside of the clinic should be determined and should be under strong control. Moreover, each silver nanomaterial should be considered as a separate agent with a potential different mode of antibacterial action.

Keywords: silver, antibiotics, resistance, sil genes, mutant, adaptant

Introduction

The overuse of antibiotics has led to the increase of bacterial resistance.¹ Currently, one of the most popular alternatives and often a supporting way, recommended as an antibacterial factor in medicine and in most industrial branches, is silver. Silver, among other metals, possesses biological activity and its high efficacy against a broad range of microorganisms (including fungi, Gram-negative and Gram-positive

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Molecular mechanisms of bacterial resistance to silver ions are well known, especially in Gram-negative bacteria.^{12,13} The first described bacterial silver resistance occurred in 1975, in Massachusetts General Hospital. Gram-negative Salmonella Typhimurium, which caused the death of several burn ward patients, was the host of the pMG101 plasmid, a 180 kb construct with specific regions involved in bacterial resistance to silver ions.^{12,14} This type of resistance was called "exogenous silver resistance" by Randall et al.¹³ The sil region covers an operon of nine sil genes, gathered in three transcriptional units (silRS, silE, and *silCFBAGP*), each controlled by a separate promoter.^{13,15} Functions of proteins encoded by sil genes were determined after comparison of their structures to homolog components of the CusCFBA and PcoE system with a chromosome-encoded efflux pump involved in the transport of copper and silver ions.^{13,16} Despite that, the function of the SilG protein remains unknown. The transcription of sil genes is carried out by the SilS/SilR two-component regulatory system, made of membrane histidine kinase and a response regulator.^{12,17} Products of *silA* (inner membrane substrate-binding transporter), silB (membrane fusion protein), and *silC* (outer membrane protein) genes form a tripartite resistance-nodulation-division (RND) efflux pump, SilCFBA - a structure spanning both membranes, responsible for ejecting silver ions out of the cell (from cytoplasm or periplasmic space).^{12,14} The protein encoded by the SilP gene is a P-type ATPase located in the inner membrane. It transports all of the toxic Ag⁺ from cytoplasm to periplasmic space, where it can be captured and delivered to SilCFBA by periplasmic chaperone - SilE.¹⁵ A similar role is performed by another protein, encoded by the silF gene. It stops the uptake of silver ions from the environment, by binding Ag⁺ in periplasmic space and passing them to the efflux pump.^{14,16} The distribution of sil genes in pMG101 plasmid is presented in Figure 1. The presence of pMG101 in bacteria is not a requirement for changing their phenotypes to silverresistant. Randall et al¹³ proved that silver ions exert selective pressure on bacterial cells treated during 6 days with silver ions (Ag⁺), causing point mutations in the bacterial (Escherichia coli) genome, e.g. in ompR or cusS genes, leading to the loss of their function. OmpR is a transcription factor of OmpC and OmpF – two outer membrane proteins, porins with a β -barrel structure, responsible for the transport of drugs and cations such as silver ions. CusS is a histidine kinase, a component of the regulatory system of chromosome-encoded efflux pump CusCFBA.^{12,17} As described by Randall et al,¹³ the examined bacteria became 6 times more resistant to silver ions. Randall et al¹³ proved that there were no changes in Staphylococcus aureus sensitivity to silver ions after 42 days of exposition to Ag^+ . Despite that, the

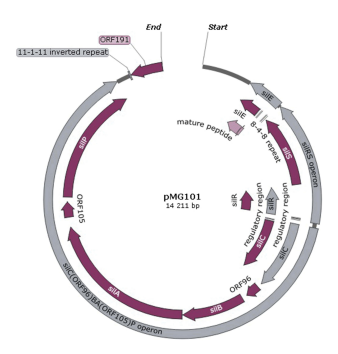


Figure 1 Map view. Distribution of sil genes in the plasmid pMG101 (created with SnapGene).

occurrence of a single *sil* gene (*sil*E) in staphylococci is possible but with an unknown function.¹⁸ Besides silver, resistance to mercury, tellurite and a few antibiotics such as streptomycin, chloramphenicol, tetracycline, and ampicillin are encoded on the pMG101 plasmid.¹²

Resistance to antibiotics is a major and growing public health problem. It is estimated that every year about 25,000 patients in Europe die due to infections caused by bacterial strains resistant to antibiotics.¹⁹ The increasing percentage of isolates of bacteria, both Gram-negative and Gram-positive, resistant to many antibiotics is indicated at the same time.²⁰

The term "multiple resistance" (MR) refers to microorganisms resistant to more than one antibiotic used in the treatment of infection.²¹ Bacteria described as "multidrug resistant" (MDR) are characterized by bacterial resistance to antibiotics that belong to 3 or more different classes.²¹ MDR strains are a group of microorganisms that are especially dangerous due to the fact that therapeutic options are limited. Data on the phenomenon of antibiotic resistance pathogens in European countries were included in the European Centre for Disease Prevention and Control (ECDC) report (November 2017) summarizing data for 2013-2016 from 30 European countries.²⁰ According to the report, the percentage of antibiotic-resistant strains is higher in the countries of southern and south-eastern Europe in comparison to the countries of northern Europe. Over 1/3 of the isolates of Klebsiella pneumoniae in Europe are resistant to one therapeutic group, but resistance to 3 or more therapeutic groups was noticed more often. Among the E. coli in European countries, a statistically significant increase in resistance was observed in the case of using third-generation cephalosporins, fluoroquinolones, and aminoglycosides. In 2013-2016, the amount of carbapenem-resistant strains of E. coli and K. pneumoniae increased in several countries. In the years 2013-2016, the European trend of methicillin-resistant S. aureus gradually decreased, although in 10 countries the percentage of such strains still remains above 25%. In Poland, since 2013, it has remained at the level of 10-25%. As a result, standard treatments become ineffective, and infections persist and may spread.¹⁹ Bacteria producing carbapenemases (usually belonging to K. pneumoniae, Pseudomonas aeruginosa, and Acinetobacterspp.) become a significant problem. The

treatment of this infection is limited to one or two antibiotics.^{22,23} The rapid spread of carbapenemase resistance (CPE) in *Enterobacteriaceae* is one of the most serious problems for the health and safety of patients. Prudent use of silver nanoformulations and widespread observance of the principles of control and prevention of infections in all health care sectors, e.g. hospitals and nursing homes, form the basis for effective interventions to prevent the selection and spread of antibiotic-resistant bacteria.^{24,25} Otherwise, Europe may soon be faced with epidemic hospital diseases caused by *Enterobacteriaceae* with extended multiple resistance (extensively drug resistant, XDR), or even strains completely resistant (pandrug resistant, PDR).²⁶

The main purpose of this study was to check the sensitivity of bacteria to silver nanoformulations after long-term exposure, to analyze the exogenous and endogenous resistance to silver, and, finally, to compare the antibiotic susceptibility of Gram-positive and Gramnegative bacteria strains (wild types and variants with changed sensitivity after repeated exposure to silver nanoformulations).

Materials And Methods Bacteria Strains

The wild types (19 strains altogether) and their selected variants (signed in the text as S1 V, S2 V, S7 V strains altogether) of Gram-negative strains (*E. coli, K. pneumoniae, Enterobacter cloacae*) and Gram-positive bacteria strains (*S. aureus*) belonging to MR or MDR bacteria were tested. Bacterial strains (wild type) were purchased from ATCC or provided by Dialab Medical Laboratory. The variants were selected after repeated exposure to silver nanoformulations (S0–S7), described below.

Chemicals

The media (Luria Broth [LB], Mueller Hinton Agar [MHA], Mueller Hinton Broth [MHB]) were purchased from Biocorp. Antibiotics were obtained from Oxoid. The Plasmid Mini Kit, Genomic Mini Kit, and DNA Ladder marker (0.1 μ g/mL. 100–1000 bp) were purchased from A&A Biotechnology. Midori Green Advance DNA Stain (Nippon Genetics), Phusion Hot Start II High Fidelity PCR Master Mix, and 50× Tris–acetate EDTA buffer (TAE, pH 8.3) were purchased

from Thermo Scientific. All of the primers were ordered in Genomed. The sequences of all used primers are summarized in Table 1 (based on the information described by Woods et al²⁷). Silver nanoformulations S1–S6, described in detail in our previous study,^{5,28} were prepared by A. Kedziora. The formulations S0 and S7 were commercially available. All silver nanoformulations are summarized in Table 2.

Long-Term Exposure Of Bacteria To Silver Nanoformulations

This task was performed with long-term exposure in gradual increases of minimal inhibitory concentration (MIC) of silver nanoformulations S0–S7 (0.5 MIC, 1 MIC, and 4 MIC). Initially, the sensitivity of bacteria to silver nanoformulations was determined via MIC measurement.²⁹ Bacteria strains were stored at $-70 \, ^{\circ}$ C before each investigation and the standard protocol was performed to check their purity and revitalization. The inoculum was spread on the agar plate (MHA),

Table I	Sequences	Of Forward And	Reverse Primers
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incubated at 37 °C for 19 h, and after that used in the experiment. In the first step, the stock of silver samples was prepared (at concentrations between 2048 µg/mL and 0.03125 µg/mL, depending on the nanoformulation sample). 100 μ L of each concentration was poured to hole. Next, the bacteria strains at the final concentration of 1.5x10⁶ cfu/mL were added and incubated at 37 °C. Minimal inhibitory concentration values were read within 16-19 h. The experiment was repeated 3 times. For 0.5 MIC and 1 MIC, broth (MHB) was used; hence, in the case of 4 MIC, an agar plate (MHA) was applied. The appropriate bacteria density (cfu/mL comparable with MIC determination) was poured into the 0.5 MIC and incubated for 24 h at 37 °C in a final volume of 10 mL. After that time, 0.5 mL of overnight culture was transferred to the new broth (containing 0.5 MIC concentration of the appropriate silver nanoformulations) and incubated under the same condition. Overnight culture of bacteria was transferred to the new broth (with 1 MIC), incubated at 37 °C forr 24 h, and the steps were repeated for 1 MIC as described for 0.5 MIC. The

sil Genes	Primers	
	Forward	Reverse
silE	AGGGGAAACGGTCTGACTTC	ATATCCATGAGCGGGTCAAC
silRS	GGCAATCGCAATCAGATTTT	GTGGAGGATACTGCGAGAGC
silCBA	CGGGAAACGCTGAAAAATTA	GTACGTTCCCAGCACCAGTT
silF	CGATATGAATGCTGCCAGTG	ATTGCCCTGCTGAATAAACG
silB	CAAAGAACAGCGCGTGATTA	GCTCAGACATTGCTGGCATA
silA	CTTGAGCATGCCAACAAGAA	CCTGCCAGTACAGGAACCAT
silP	CCTGGGTTTACAGCGTCATT	ATGGCACCTGAGGTTTGTTC

Note: Data from Woods EJ, Cochrane CA, Percival SL. Prevalence of silver resistance genes in bacteria isolated from human and horse wounds. Vet Microbiol. 2009;138:325-329.²⁷

Table 2 Silver Nanoformulations	, Which	The Bacteria	Were Treated Wit	ch
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Silver Nanoformulations	Details	References/Origin
SO	Silver nanoparticle powder without carrier and stabilizers; silver size <100 nm	Commercially available
SI	Silver ions immobilized on amorphous TiO_2 (TiO_2/Ag^+)	28
S2	Silver nanoparticles immobilized on amorphous TiO ₂ (TiO ₂ /Ag ⁰), silver size 20 nm	28
S3	Silver ions immobilized on crystalline TiO ₂ (TiO _{2a} /Ag ⁺)	28
S4	Silver nanoparticles immobilized on crystalline TiO ₂ (TiO _{2a} /Ag ⁰), silver size ≤ 10 nm	28
S5	Silver nanoparticles immobilized on amorphous TiO₂ (TiO₂:Ag ⁰), silver size ≤5 nm	5
S6	Silver nanoparticles immobilized on crystalline TiO₂ (TiO₂a:Ag⁰), silver size ≤5 nm	5
S7	Aqueous dispersion of silver nanoparticles; stabilizers: trace amount of Tween and	Commercially available
	polyethyleneimine, silver size 20 nm	

Notes: Forms had different physicochemical properties: lack or presence of carrier or stabilizers, carrier size, silver size, surface area, and chemical compounds.

	E. cloacae 233 S7 V	n/a 8 n/a 5 n/a 512 n/a n/a 8 n/a n/a n/a 8 n/a n/a n/a 16 n/a 1 and in Table 2; S0, silver 3 3 3
	Е. сіоасае 233	8 512 8 16 0.5 ble 2; S
	K. pneumoniae 268 S7 V	n/a n/a n/a 4 d in Ta
	K. pneumoniae 268 wt	8 256 16 16 0.5 0.5 ²⁸ an
	K. pneumoniae 626 S7 V	n/a n/a n/a 64 64 iora et viously ^{5,}
	K. pneumoniae 626 S2 V	n/a n/a n/a 64 n/a y Kędz
	K. pneumoniae 626 SI V	n/a n/a 8 n/a n/a cribed l
	K. pneumoniae 626 wt	4 n/a n/a 8 n/a 8 n/a 8 n/a 8 n/a 8 n/a 8 n/a 1/a 1/a <th1< th=""></th1<>
	K. pneumoniae ATCC 4352 S7 V	n/a n/a n/a n/a l n/a n/a n/a
	K. pneumoniae ATCC 4352 S2 V	8 n/a n/a n/a icochem
	K. pneumoniae ATCC 4352 SI V	n/a n/a 16 n/a n/a ss, phys
	K. pneumoniae ATCC 4352 wt	8 256 0.5 4* 1 Ifference mpound
	E. coli 343 S7V	n/a n/a n/a n/a 4 mical di
sr	E. coli 343 wt	8 256 8 16 2 sicoche
Strair	E. coli 408 S7V	n/a n/a n/a n/a 2 vith phy centrati
Icteria	E. coli 408 S2 V	n/a n/a n/a 16 n/a to con to con
cive Ba	E. coli 408 wt	8 256 8 4 1 0 silver
-Negat	E. coli 555 S7 V	n/a n/a n/a n/a 2 7, kind ce. *Co
Gram.	E. coli 555 S2 V	n/a n/a n/a 8 n/a 1, S2, S ¹
ested (E. coli 555 w t	4 256 0.5 4* 1 1 ctions; S tions; S
inst Te	E. coli 475 S7 V	n/a n/a n/a 4 MG101
ıs Aga	E. coli 475 S2 V	n/a n/a n/a 32 n/a er nano
ulation	E. coli 475 w t	4 256 0.5 4* 1 vith silve
oform	E. coli ATCC 11229 S7 V	n/a n/a n/a 4 CPositi
. Nano	E. coli ATCC 11229 S2 V	n/a n/a n/a 32 32 n/a n/a able 2).
Silver	E. coli ATCC 11229 wt	4 256 0.5 4* 4* 1 Iong-ter long-ter ain: n/a.
Different	E. coli J33	256 >2048 256 16 16 cted after ercially ava did type stri
Table 3 MIC Of Different Silver Nanoformulations Against Tested Gram-Negative Bacteria Strains	Silver nanoformulations strains	$ MIC \text{ Ag}^{\dagger} \left[\text{ Jg}/\text{Im} \right] \ 256 44 n/a n/a 16 n/a n/a $
F		 z e Z

Bacteria strains											
Silver nanoformulations S. aureus ATCC 6	V 72 8526 DDTA _{2U9} via. 2	S. aureus 173	V 72 EVI eureus 173 S7 V	S. aureus 298	S. aureus 298 S7 V	S. aureus 187	A29M 045 suprus .2	V 02 A2AM 045 2usine .2	V 72 A29M 045 sugare S.	S. aureus 2	S. aureus 2 MRSA S7 V
MIC Ag ⁺ [µg/mL] 8	n/a	8	n/a	2	n/a	4	32	32	n/a	32	n/a
MIC S0 [µg/mL] 1600	n/a	1600	n/a	1600	n/a	1600	2048	4096	n/a	2048	n/a
MIC SI [µg/mL] 0.5	n/a	0.5	n/a	0.5	n/a	0.05	16	n/a	n/a	8	n/a
MIC S2 [µg/mL] 4*	n/a	4 *	n/a	4 *	n/a	4 *	I6	16	n/a	16	n/a
MIC S7 [µg/mL] I	2	_	2	_	2	_	_	n/a	2	_	2

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population that adapted to the increasing concentration of silver nanoformulations grew on the agar plate (4 MIC). The MIC value was then determined again as described above and compared with that before longterm exposure.³⁰

Analysis Of Whole Genome Sequences

Whole genome DNA was isolated with the Genomic Mini Kit according to the original instruction from 2 mL of overnight LB cultures. The product purity and concentration were verified with a Nano Photometer (Implen). Genomic libraries were prepared according to the NEBNext DNA Library Prep Master Mix Set for Illumina protocol, and sequencing (NGS) was performed at Genomed (Warsaw, Poland) using the Illumina MiSeq platform. NGS reads were trimmed with Cutadapt 1.9.1.31 De-novo genome assembly was performed with SPAdes 3.13.0,³² their quality was evaluated with QUASTt 5.0.2,³³ and resulting contigs were rearranged with progressiveMauve in Mauve 2.4.0^{34,35} using the genome of E. coli K-12 substr. MG1655 (NC 000913.3) or K. pneumoniae subsp. pneumoniae HS11286 (NC 016845.1) as a reference. Newly assembled genomes were annotated with prokka 1.12.36 Genomic mutations in adapted strains were identified and characterized with snippy 4.3.8.37 The pre-processed reads and assembled genomes have been deposited in the NCBI SRA and Genome databases accordingly under the following accession numbers: SRR9733699, SRR9733700, SRR 9733697, SRR9733698, SRR9733703, SRR9733704, SRR 9733701, SRR9733702, SRR9733705, SRR9733706, SRR9 733707 and VLTC00000000, VLTB00000000, VLTA000 00000, VLSZ00000000, VLSY00000000, VLSX00000 000, VLSW00000000, VLSV00000000, VLSU00000000, VLST0000000, VLSS00000000.

Detection Of Sil Genes

To verify the exogenous silver resistance in all tested bacteria strains, the prevalence of sil genes located on pMG101 plasmid was determined in the following steps.

Plasmid Isolation

Abbreviations: wt, wild type strain; n/a, not applicable

Plasmid pMG101 was isolated with the Plasmid Mini Kit according to the original instruction. The product quantity of this extraction was confirmed with the Nano Photometer (Implen). Plasmid DNA was isolated from all of the tested strains (described above), and the one from the E. coli J53 strain was used as a positive

								Bac	teria s	strains						
	E. coli J53	<i>E. coli</i> ATCC 11229	E. coli 475	E. coli 555	E. coli 408	E. coli 343	K. pneumoniae ATCC 4352	K. pneumoniae 626	K. pneumoniae 268	E. cloacae 233	S. aureus ATCC 6538	S. aureus 173	S. aureus 298	S. aureus 187	S. aureus 340 MRSA	S. aureus 2
So	n/a	ı	ı	ı	ı	ı	ı	ı	ı	ı	Т	ı	ı	ı	+	1
S1	n/a	ı	I	ı	I	I	+	+	I	ı	I	ı	I	I	I	ı
S2	n/a	+	÷	+	+	ı	+	+	ı	ı	I	ı	ı	I	I	ı
S3	n/a	I	ı	ı	ı	1	1	ı	1	ı	I	ı	ı	I	I	ı
S4	n/a	I	ı	ı	ı	1	ı	ı	ı	ı	I	ı	1	ı	ı	1
S5	n/a	I	I	ı	ı	ı	ı	ı	I	ı	I	ı	ı	ı	ı	1
S6	n/a	I	ı	ı	ı	1	1	ı	ı	ı	I	ı	1	ı	ı	1
S7	n/a	+	+	+	+	+	ı	+	+	+	+	+	+	I	÷	+

Table 5 The Changes Of Bacterial Sensitivit	ty To Silver Nanoformulations After Their Repeated Exposure
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Notes: "+", increases of MIC after 6 days of exposition to appropriate silver nanoformulations; "-", no changes of MIC noticed after 6 days of exposition to appropriate silver nanoformulations, no variants obtained. Variants with changed sensitivity to silver formulations (S0–S7). Abbreviation: n/a, not applicable.

control. All bacteria strains grew on Luria Broth before DNA extraction.

Primer Preparation

In the first step, lyophilized primers were centrifuged (12,500 rpm, 2 min); next, the required volume of water (MQ) was poured to the final concentration of 100 μ M. The primers were vortexed and left for 10 min at room temperature. Then, they were diluted 10 times and stored at -20 °C.

PCR Mixture

The following reagents were mixed for every reaction: 5 μ L Phusion Hot Start II High Fidelity PCR Master Mix, 0.5 μ L forward primer, 0.5 μ L reverse primer, 1 μ L plasmid, and the required volume of MQ. The final volume of all reactions was 10 μ L. The positive (with plasmid from *E. coli* J53) and negative (sample without any of plasmid) controls were performed.

PCR Conditions

The following steps were established for the best final products of PCR: 30 sec at 98 °C (initial denaturation) followed by 30 cycles of 98 °C for 5 sec (denaturation), 60 °C for 10 sec (annealing), and 72 °C for 12 sec (elongation); and 72 °C for 10 min (final elongation). PCR was carried out on a T100TM Thermal Cycler (BioRad).

DNA Electrophoresis

5 μ L of each sample and 1 μ L of heavy buffer (30% glycerol stained bromophenol blue) were loaded to a 2% agarose Tris–acetate–EDTA (TAE) gel containing 0.002%

Table 6 Number And Dist	Table 6 Number And Distribution Of Genomic Mutations Acquired By Selected Test Variants (V) Due To Long-Term Exposure For Different Silver Nanoformulations	tions Acq	luired	By Se	lected Te	st Varia	nts (V)	Due	To Long-	Ferm Ex	posure	For L	Different	Silver Nan	oformulatic	suc
Silver Nanoformulations Variants (V)	Variants (V)	Genomic Region	ic Reg	ion										Genes W	Genes With Mutations	su
		CDS				RNA				Intergenic	nic					
		SNP	Del	lns	Comp	SNP	Del	lns	Comp	SNP	Del	lns	Comp	Del Ins Comp In Total	Nonconservative	rvative
															Number	Examples
SI	K. pneumoniae ATCC 4352	20,375	67	26	1663	26	2	2	3	2680	87	88	476	19,008	829	Inc. cusS tamB
S2	E. coli ATCC 11229	15,639	5	9	2429	I	I	2	m	1331	34	29	298	16,477	388	Inc. cusS
SI	K. pneumoniae 626	2	I	I	Ι	I	I	I	I	I	ı	I		2	_	Only cusS
S2	K. pneumoniae 626	2	I	I	Ι	I	I	I	I	2	1	I	e	2	_	Only cusS
S2	K. pneumoniae ATCC 4352	_	I	I	I	I	I	I	I	I	I	1	1	_	_	Only tamB
S7	K. pneumoniae ATCC 4352	I	I	I	Ι	I	I	I	I	I	ı	I		I	I	I
S7	E. coli ATCC 11229	з	I	I	I	I	I	I	I	I	1	I	1	2	I	I
S7	K. pneumoniae 626	I	I	I	I	I	I	I	I	2	I	1	e	I	I	I
Notes: 2–53 nt in length, average 4 nt.	4 nt.										1					

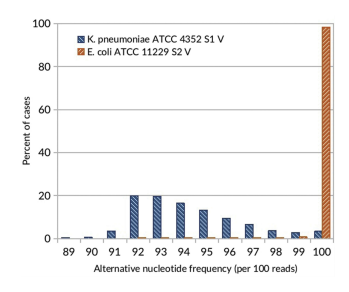


Figure 2 A histogram showing mutation frequency counts in genomic sequencing of *Escherichia coli* ATCC 11229 S2 V and *Klebsiella pneumoniae* ATCC 4352 S1 V strains.

Midori Green. Electrophoresis was carried out during 0.5 h at 7.5 V/cm. A 5 μ L DNA marker was applied on the gel. Images of the gels were carried out on a Gel Doc XR⁺ (BioRad).

Antibiotic Susceptibility Testing

Sensitivity of the tested bacterial strains to antibiotics was determined with disc diffusion methods. The criteria for selection of antimicrobials were based on the EUCAST recommendation.³⁸ Bacteria (0.5 McFarland) were inoculated and antibiotic discs were placed on the MHA plate. Plates were incubated at 37 °C during 18 h, followed by zone diameter measurement and breakpoint estimation.

Results And Discussion

Abbreviations: CDS, protein coding sequences; SNP; single nucleotide polymorphism; Del, deletion; Ins, insertion; Comp. complex

The changes of bacterial sensitivity to silver nanoformulations were determined after bacteria strains' exposition to increasing concentrations of silver nanoformulations. After comparison of the primary (in wild type strains, wt) and secondary (among variants, V) MIC values, the changes of bacterial sensitivity were determined and the selection of variants was possible. The obtained MIC values are described in Tables 3 and 4. The susceptibility of bacteria depended on the kind of silver nanoformulations (their size, shape, surface area, compounds, etc.) and bacterial strains. Among some of the tested bacterial strains, the changes of sensitivity to certain silver nanoformulations were noticed. It is interesting that almost all strains

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	E. coli K-12 PFM2 wt ⁴³	<i>E. coli</i> K-12 PFM2 mutl. ⁴³	<i>E.coli</i> ATCC 11229 S2 V	K. pneumoniae ATCC 4352 S1 V	K. pneumoniae 626 S1 V	K. pneumoniae 26 S2 V	K. pneumoniae ATCC 4352 S2 V	E. coli ATCC 11229 S7 V	K. pneumoniae 626 S7 V
Substitutions in total	233	1 625	17 001	23 081	2	4	1	3	2
Type of substitution									
Transitions									
A:T>G:C G:C>A:T		49 82 447	1 5 988 6 68 ⁴		1	2 2		- 1	1 1
Transversions									
A:T>T:A A:T>C:G G:C>T:A G:C>C:G		17 14 38 10 30 10 17 3	1 143 1 192 1 220 777	2 1 682	- 1 -	-	1 - -	1 - 1	
Consequences of substitut	-	5		1 040	_			-	_
Position	0115								
Noncoding Coding Within coding sequences		54 213 179 1412			- 2	2 2	- 1	- 3	2
Synonymous		55 482	13 73	2 16 727	1	1	1	3	_
Nonsynonymous		124 930	1 907	7 3 648	1	1	_	_	_
Amino acid changes					1	1	1		1
Conservative		57 644			1	1	-	3	-
Nonconservative		67 286	477	1 077	-	1	10	-	

Table 7 Substitution C	Jounts From	Whole Genome	e Sequencing As	Divided For	Transitions /	And Transversions,	Followed By
Distribution Of Substitut	tions In Codin	ng Regions And F	ossible Impact O	n Coded Prot	ein Structure		

Notes: Green and blue marks illustrate relative abundance of particular substitution type.

changed their susceptibility level to S7, 4 of 5 E. coli changed sensitivity to S2, 2 of 3 K. pneumoniae changed sensitivity to S1 and S2, and only 1 of 6 S. aureus changed the sensitivity to S0. All of the selected variants are summarized in Table 5. To verify the reasons for changing the variants' sensitivity, whole genome sequences were analyzed. DNA from 11 of the tested wt strains and their phenotype variants (V) was isolated: K. pneumoniae ATCC 4352 wt, K. pneumoniae ATCC 4352 S1 V, K. pneumoniae ATCC 4352 S2 V, K. pneumoniae ATCC 4352 S7 V, K. pneumoniae 626 wt, K. pneumoniae 626 S1 V, K. pneumoniae 626 S2 V, K. pneumoniae 626 S7 V, E. coli ATCC 11229 wt, E. coli ATCC 11229 S2 V, and E. coli ATCC 11229 S7 V. The above strains were selected on the basis of the obtained results described in Table 5. For the analysis, mutations were identified in protein coding sequences (CDS), RNA coding sequences, and intergenic regions. All mutations recorded in the above genomic regions were divided into nucleotide substitutions, insertions, deletions, and other complex mutations (Table 6). The most mutations (mainly SNP) were noticed among K. pneumoniae ATCC 4352 and E. coli ATCC 11229 strains treated with S1 and S2, respectively. Definitely fewer genomic changes were indicated among K. pneumoniae 626 treated with S1 and S2, and K. pneumoniae ATCC 4352 treated with S2. In general, all mutations can be divided into conservative and nonconservative. Mutation is considered as conservative when it does not introduce any change in primary protein structure or the introduced amino acid change has score >0 in BLOSUM62.³⁹ Otherwise, mutation is considered as nonconservative.⁴⁰ Counts of genes harboring at least one mutation (including conservative and nonconservative) are listed in Table 6. In the case of the strains with the highest number of mutations (K. pneumoniae ATCC 4352 S1 V and E. coli ATCC 11229 S2 V), more than 1200 genes were identified with nonconservative mutations (only a few examples are shown in Table 6), while among the strains K. pneumoniae 626 S1 and S2 V and K. pneumoniae ATCC

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Bacteria Strains	sil Genes						
	silE	silRS	silCBA	silF	silB	silA	silP
E. coli J53	+	+	+	+	+	+	+
E. coli ATCC 11229	+	+	+	+	+	-	-
E. coli 475	-	-	-	-	-	-	-
E. coli 555	-	-	-	-	-	-	-
E. coli 408	-	-	-	-	-	-	-
E. coli 343	-	-	-	-	-	-	-
K. pneumoniae ATCC 4352	-	-	-	-	-	-	-
K. pneumoniae 626	-	-	-	-	+	-	-
K. pneumoniae 268	-	-	-	-	-	-	-
E. cloacae 233	-	-	-	-	-	-	-

Notes: +, presence of gene; -, lack of gene.

4352 S2 V only one gene was identified, cusS and tamB appropriate. It is worth underlining that the same genes were noticed in cases of strains with a huge number of mutations. The obtained results can be divided into three clusters, where each one has a distinctive feature: a) high number of mutations (*E. coli* ATCC 11229 S2 V, *K. pneumoniae* ATCC 4352 S1 V), b) single gene mutations (*K. pneumoniae* 626 S1 and S2 V, and *K. pneumoniae* ATCC 11229 S7 V and, *K. pneumoniae* ATCC 4352 S7 V and 626 S7 V) (Table 6). The mutation frequency counts in genomic sequencing of *E. coli* ATCC 11229 S2 V and *K. pneumoniae* ATCC 4352 S1 V strains are shown in Figure 2. As indicated by these data, all recorded mutations occurred with a frequency between 82 and 100 per 100 reads, suggesting some level of heterogeneity gained in silver-treated

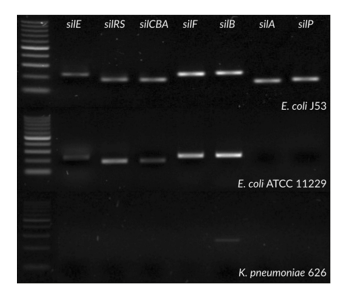


Figure 3 Electropherograms of Escherichia coli ATCC 11229 wt and Klebsiella pneumoniae 626 wt sil genes.

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selective pressure depending on the individual features of bacterial strains, in contrast to S7. To explain the mutation rates among obtained variants after S0-S7 exposition, the substitution counts from whole genome sequencing followed by the distribution of substitutions in coding regions and possible impact on coded protein structure are described in Table 7. Since the two variants (E. coli ATCC 11229 S2 V and K. pneumoniae ATCC 4352 S1 V) gained an extraordinarily high number of mutations, they may be premised as mutator strains.⁴¹ This observation can be explained by a possible decrease of DNA repair efficiency in obtained variants, resulting from nonconservative mutations in genes coding proteins involved in mismatchrepair (MMR): mutL (Pro389Ser, Leu418Pro) in E. coli ATCC 11229 S2 V, and mutL (Pro371Thr) and mutM (Ala148Glu) in K. pneumoniae ATCC 4352 S1 V. As MMR deficiency promotes not only quantitative, but also qualitative changes in the mutational profile, substitution rates from this experiment are compared with corresponding data obtained by Lee et al.⁴⁰ In brief, the wt strain gains transitions and transversions with comparable ratios (56% vs 44%), with a slight domination of G:C>A:T transitions. In contrast, the MMR defective strain (mutL) gains mostly transitions (over 90%) with clear dominance of A:T>G: C transitions (70%). The two mutator-like variants from this experiment, carrying spontaneous mutation in *mutL*, do not fully fall into any of these categories, although the transition ratio (73–75%) is notably higher than the transversion ratio

Table 9 Antibiotics Susceptibility Of Gram-Negative Bacteria Strains

Bacteria Strains		Am	АМС	СХМ	стх	CAZ	ATM	AN	CIP	sхт	Imp	Mem	TZF
E. coli J53 ^C		-	R	R	R	s	R	-	s	S	s	s	R
E. coli ATCC 11229 wt		s	s	s	s	s	s	S	s	s	s	s	s
E. coli ATCC 11229 S2 V		s	s	s	s	s	s	s	s	S	s	S	s
E. coli ATCC 11229 S7 V		s	s	s	s	s	s	s	s	S	s	s	s
E. coli 475 wt	MR	R	s	s	s	s	s	S	S	s	s	s	s
E. coli 475 S2 V		R	s	s	s	s	s	S	R	s	s	s	s
E. coli 475 S7 V		S	s	s	s	s	S	s	s	S	s	S	s
E. coli 555 wt		R	R	R	s	s	s	S	s	s	s	s	s
E. coli 555 S2 V		R	R	S	s	s	s	S	s	s	s	s	s
E. coli 555 S7 V		R	R	S	s	s	s	s	s	S	s	s	s
E. coli 574 wt	MDR	R	R	R	s	s	s	S	s	S	s	s	s
E. coli 574 S7 V		S	S	S	s	s	s	S	s	s	s	s	s
E. coli 408 wt		R	R	R	R	s	s	s	s	R	s	s	s
E. coli 408 S2 V		R	R	R	R	s	s	S	s	R	s	s	s
E. coli 408 S7 V		S	S	S	S	s	S	S	s	S	s	S	s
E. coli 343 ESBL wt		R	R	R	R	R	R	R	R	R	s	S	s
E. coli 343 ESBL S7 V		S	R	R	R	S	S	S	R	R	s	s	s
K. pneumoniae ATCC 4352 wt		R	R	R	R	R	R	s	s	S	s	S	s
K. pneumoniae ATCC 4352 S2 V		R	R	R	R	R	R	S	s	S	s	S	s
K. pneumoniae ATCC 4352 S7 V		S	S	S	S	S	S	s	s	s	s	s	s
K. pneumoniae 626 wt		R	R	R	R	R	R	S	s	S	s	S	s
K. pneumoniae 626 S2 V		R	R	R	R	R	R	s	s	S	s	S	s
K. pneumoniae 626 S7 V		R	R	R	R	R	S	s	s	S	s	S	s
K. pneumoniae 268ESBL wt		R	R	R	R	R	R	S	R	S	s	S	s
K. pneumoniae 268ESBL S7 V	1	S	R	R	R	S	S	s	R	s	s	s	s
E. cloacae 233 wt		R	R	R	R	R	R	S	S	R	s	s	s
E. cloacae 233 S7 V		S	R	R	R	S	S	s	R	R	s	s	s

Notes: V, variants selected after long-term treatment with silver nanoformulations; S2, S7, etc., kind of silver form with physicochemical differences. ^CPositive control of pMG101 plasmid presence; green marks, increasing the susceptibility of variants to antibiotic; red marks, decreasing the susceptibility of variants to antibiotic; gray marks, no changes, no differences between wild type and variants.¹ No variants obtained.

Abbreviations: wt, wild type strain; S, susceptible; R, resistant; Am, ampicillin; AMC, amoxicillin-clavulanic acid; CXM, cefuroxime; CTX, cefotaxime; CAZ, ceftazidime; AN, amikacin; ATM, aztreonamum; CIP, ciprofloxacin; Imp, imipenem; Mem, meropenem, TZP, piperacillin-tazobactam; SXT, trimethoprim-sulfamethoxazole.

(25–27%), what is a resemblance to the *mutL* strain, where A: T>G:C and G:C>A:T transition ratios are close to equal. Considering the effect of substitution, in both WT and *mutL* strains, the nonsynonymous mutation ratio outweighed the synonymous mutation ratio. Interestingly, in silver-resistant

variants the trend is opposite and most substitutions do not affect coded proteins. Therefore, the mutator phenotype in *E. coli* ATCC 11229 S2 and *K. pneumoniae* ATCC 4352 cannot be explained with a single *mutL* dysfunction, and may be the result of more complex changes and specific selective pressure

susceptibility of variants to antibiotic; red marks, decreasing the susceptibility of variants to antibiotic; gray marks, no changes, no differences between wild type and variants. **Abbreviations**: wt, wild type strain; S, susceptible; R, resistant; AMC, amoxicillin-clavulanic acid; CXM, cefuroxime; CAZ, ceftazidime; AN, amikacin; Ge, gentamicin; CIP, ciprofloxacin; Imp, imipenem; Mem, meropenem; SXT, trimethoprim-sulfamethoxazole; P, penicillin; E, erythromycin; CC, clindamycin; FOX, cefoxitin; Link, lincomycin; Tet, tetracycline; C, chloramphenicol.

Bacteria Strains	٩	AMC	СХМ	СТХ	CAZ	Ge	AN	сњ	sхт	d m	Mem	ш	U U	Va Fox	x Link	< Tet	υ
S. aureus ATCC 6538	R	S	S	S	S	s	S	R	R	s	S	R	R	s s	Я	R	R
S. aureus ATCC 6538 S7 V	S	S	S	S	S	S	S	S	S	S	S	S	S S	s s	S	S	R
S. aureus 173 wt	R	R	R	R	R	R	R	R	R	R	R	R	R	S R	R	S	S
S. aureus 173 S7 V	S	S	S	S	S	S	S	S	S	S	S	S	s s	s s	S	S	S
S. aureus 298 wt	R	S	S	S	S	S	s	~	s	S	S	۲	R	s s	R	s	Я
S. aureus 298 S7 V	S	S	S	S	S	S	s	S	s	S	S	s	s, s	s s	S	S	S
S. aureus 187 MLSB MRSA wt	R	R	R	R	R	R	Я	Я	R	Я	R	R	R	s R	R	R	S
S. aureus 340 MRSA wt	R	R	R	R	R	R	S	S	R	R	R	S	S	S R	S	S	R
S. aureus 340 MRSA S0 V	Я	R	R	R	R	R	۲	s	S	R	Я	۲	R	S R	۲	۲	S
S. aureus 340 MRSA S7 V	S	S	S	S	S	S	S	S	S	S	S	S	s s	s s	S	S	S
S. aureus 2 MRSA wt	R	R	R	R	S	R	R	R	R	R	R	R	R	S R	R	S	R
S. aureus 2 MRSA S7 V	S	S	S	S	S	S	S	S	S	S	S	S	s	s s	S	S	S
Notes: V, variants selected after long-term treatment with silver nanoformulations;	ns; S2,	S7, etc.,	kind of si	kind of silver form with physicochemical differences.	with phy	sicochen	nical diff		^C Positive control of $pMG101$ plasmid presence;	control	of pMGI	01 plas	smid pre	sence; gr	green marks, increasing the	s, increa	sing the

Table 10 Antibiotics Susceptibility Of Gram-Positive Bacteria Strains

of silver nanoformulations. Other examined variants do not harbor mutations in any known gene related to DNA repair, consistently the mutator phenotype is missing in them. Table 7 illustrates the relative abundance of particular substitution types within the sample. Genotoxicity of silver nanoparticles toward Gram-positive and Gram-negative bacteria was confirmed.⁴² Panacek et al⁴³ showed that Gram-negative bacteria (including E. coli strains) can develop resistance to silver nanoparticles after repeated exposure but without any genetic changes. It resulted from phenotypic changes involving inhibition of the flagellum protein production. Panacek et al⁴³ confirmed that strains which changed sensitivity to silver nanoparticles did not change sensitivity to silver ions. Those results suggest different response of bacteria cells to various types of silver but is the opposite of studies performed by Randall et al¹³ that proved the genomic changes in *E. coli* strains after 6 days of exposure to a subinhibitory concentration of silver ions.

Among all of the bacteria strains tested by us, both Gramnegative and Gram-positive, the presence of *sil* genes, located on the plasmid pMG101, was checked. We indicated that 2 strains possess some *sil* genes: *silE*, *silRS*, *silCBA*, *silF*, and *silB*were present in *E. coli* ATCC 11229 and *silB*in *K. pneumoniae* 626 (Table 8). It is worth underlining that those strains showed high sensitivity to silver ions and tested silver nanoformulations (excluding S0) (Table 3). Probably, this results from incompleteness of the pMG101 plasmid. In most of these strains the lack of all checked *sil* genes was confirmed. All results were visualized with electrophoresis (Figure 3).

Finally, the antibiotic susceptibility of variants was checked. Results are summarized in Tables 9 and 10. Results indicated that 4 variants (E. coli 475 S2 V and E. cloacae 233 S7 Vamong Gram-negative bacteria and S. aureus ATCC 6538 S7 V and S. aureus 340 S7 V among Gram-positive bacteria) became more resistant to antibiotic (ciprofloxacin in the case of Gram-negative bacteria and few classes in Gram-positive strains) (Tables 9 and 10). All of them were obtained after exposition to different kinds of silver nanoformulations: S2 and S7. Remaining Gram-negative and Gram-positive variants maintained the level of antibiotic susceptibility or became more sensitive to some of the antibiotic classes. Almost all of the S7 variants became more sensitive to certain classes of antibiotics. One of the possible mechanisms of bacterial resistance to antibiotics is the inhibition of antibiotic penetration into the bacteria cell.44-48 In this case it may be related to higher envelope permeability after silver nanoformulation treatment. On the other hand, Barras et al⁴⁹ reviewed that incorporation of silver into antibiotics can increase their antibacterial activity. Anuj et al⁵⁰ confirmed that cationic particles of nanosilver disturb the membrane integrity in bacteria strains, increasing their permeability in E. coli for linezolid, its higher intracellular concentration, and better antibacterial efficacy. A similar study was performed by Kaur et al⁵¹ who conjugated vancomycin with silver nanoparticles and enhanced antibacterial activity against both classes of bacteria: Gram-positive S. aureus and Gram-negative E. coli. However, Mühling et al⁵² showed that environmental exposure of resistant bacteria to silver nanoparticles did not increase antibiotic resistance in naturally occurring strains. They proved that there is no interaction between the antibiotics and Ag. As we reviewed,⁹ the interaction of bacteria with silver nanoformulations and consequences depends on the physicochemical properties of silver. The size, shape, biodiversity, and active surface determine their mode of action and the cell answer.

Conclusion

Silver nanoformulations with different physicochemical properties exert selective pressure on the bacterial strain population. Application of silver in the case of MR or MDR bacterial infection may cause different bacterial responses to antibiotics. Based on our results and the analyzed literature we may conclude that the response of bacteria cells to silver nanoparticles depends on the physicochemical properties of the nanoformulations (such as size, shape, charge, surface area, compounds) and individual features of bacteria strains (such as structural compounds and metabolism). The incorporation of silver into industrial products should be under strong control. Moreover, each silver nanomaterial should be considered as a separate agent with a potential different mode of antibacterial action.

Highlights

- 1. Silver nanoformulations, in dependence on physicochemical properties, can exert selective pressure on the bacterial cells, decreasing the sensitivity to silver.
- 2. Phenotypical changes, as a consequence of long-term exposure to silver nanoformulations, can change their sensitivity to antibiotics, including increasing bacterial susceptibility to some classes of antibiotics.
- Silver nanoformulations may cause mutational or phenotypical changes in dependence on physicochemical properties.
- 4. Development of bacterial resistance to silver nanoformulations depends on their physicochemical

properties and individual features of the bacteria cell (e.g. cell structure, cell compounds, and metabolism)

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