Bactericidal activity of alpha-bromocinnamaldehyde against persisters in *Escherichia coli*

Qingshan Shen¹*, Wei Zhou¹*, Liangbin Hu¹*, Yonghua Qi², Hongmei Ning², Jian Chen³, Haizhen Mo¹*

¹ Department of Food Science, Henan Institute of Science and Technology, Xinxiang, China, ² Department of Animal Science, Henan Institute of Science and Technology, Xinxiang, China, ³ Institute of Food Quality and Safety, Jiangsu Academy of Agricultural Science, Nanjing, China

* These authors contributed equally to this work.
  ☉ huib973@163.com (LH); mohz@163.com (HM)

Abstract

Persisters are tolerant to multiple antibiotics, and widely distributed in bacteria, fungi, parasites, and even cancerous human cell populations, leading to recurrent infections and relapse after therapy. In this study, we investigated the potential of cinnamaldehyde and its derivatives to eradicate persisters in *Escherichia coli*. The results showed that 200 μg/ml of alpha-bromocinnamaldehyde (Br-CA) was capable of killing all *E. coli* cells during the exponential phase. Considering the heterogeneous nature of persisters, multiple types of persisters were induced and exposed to Br-CA. Our results indicated that no cells in the ppGpp-overproducing strain or TisB-overexpressing strain survived the treatment of Br-CA although considerable amounts of persisters to ampicillin (Amp) and ciprofloxacin (Cip) were induced. Chemical induction by carbonyl cyanide m-chlorophenylhydrazone (CCCP) led to the formation of more than 10% persister to Amp and Cip in the entire population, and Br-CA still completely eradicated them. In addition, the cells in the stationary phase, which are usually highly recalcitrant to antibiotics treatment, were also completely eradicated by 400 μg/ml of Br-CA. Further studies showed that neither thiourea (hydroxyl-radical scavenger) nor DPTA (Fe³⁺ chelator to block the hydroxyl-radical) affected the bactericidal efficiency of the Br-CA to kill *E. coli*, indicating a ROS-independent bactericidal mechanism. Taken together, we concluded that Br-CA compound has a novel bactericidal mechanism and the potential to mitigate antibiotics resistance crisis.

Introduction

The current antibiotic resistance crisis arising from the large-scale use of antibiotics has led to increased pressure to prioritize strategies to tackle the issue in clinical practice [1]. The need for new antibiotics is more pressing due to the presence of persister cells, which are defined as a small fraction of a bacterial population displaying non-hereditary tolerance to high doses of bactericidal antibiotics [2, 3]. Persistence has been widely described in bacteria, fungi,
parasites, and even cancerous human cell populations, which ultimately led to recurrent infections and relapses after therapy [4]. A plethora of environmental signals such as nutrient limitation, pretreatment by sub-inhibitory concentrations of antibiotics, oxidative stress, heat shock, or DNA-damaging agents, can all increase persister levels, which poses a great challenge to antibiotic therapy as well [5–9].

Ubiquitous bacterial stress alarmone ppGpp is widely accepted as an emerging central regulator of persistence both in stochastically and environmentally induced persistence through its effectors, toxin-antitoxin modules [9]. In addition, induction of the SOS response dramatically increases persistence to fluoroquinolones, which is dependent on the DNA damage-inducible TisB toxin [6]. Both pathways are supported by the fact that rifampicin (Rifa, halting transcription), tetracycline (Tetra, halting translation), and carbonyl cyanide m-chlorophenylhydrazone (CCCP, halting ATP synthesis), which mimic type II and TisB-like toxins, respectively, could dramatically increase persistence [10]. Based on the known persister characteristics, several approaches, e.g., alteration of membranes via Trp/Arg-containing antimicrobial peptides [11], activation of ClpP-mediated self-digestion via Rifa and ADEP4 [12], and conversion of persisters to non-persisters via cis-2-decenolic acid [13] have been proposed. The discovery that some individual compound such as mitomycin C [14], boromycin [15], and brominated furanones [16] had the potential to eradicate persisters is also very promising [11]. Considering the diverse and variable nature of persisters [17], more drugs need to be developed based on the screening of extensive compound resources.

Ancient medical treatments against microbial infection mainly depended on herbal plant resources [18]. In particular, essential oils have frequently been found to display wide-spectrum antimicrobial activity. Cinnamaldehyde (CA), a kind of essential oil, has potent antimicrobial activity against bacteria and fungi [19, 20]. Recently, CA has been indicated to dramatically inhibit the production of biofilm in many microbes [21–23]. It has been generally accepted that persister cells are responsible for the relapse of biofilm infections [24, 25]. Therefore, we hypothesized that novel persister eradicators could be found from CA derivatives. In this study, we determined the activity of CA and its eight derivatives against persisters in *E. coli*, and found that alpha-bromocinnamaldehyde (Br-CA) had great potential to eradicate persisters.

### Materials and methods

#### Strains, culture conditions, and chemicals

The *E. coli* strains used are listed in Table 1. All experiments were conducted at 37°C in Luria-Bertani (LB) medium shaking at 220 rpm (liquid cultures) [26]. Rifa, Tetra, CCCP, ciprofloxacin (Cipro), and ampicillin (Amp) were purchased from Sigma-Aldrich. CA and its derivatives (Table 1) were synthesized and purified by Shanghai Fisher Scientific Co., Ltd.

| Strains and plasmids | Description | Source or reference |
|----------------------|-------------|---------------------|
| **E. coli** K-12 Strains |  |  |
| MG1655 | F⁻ λ⁻ ilvG rfb-50 rph-1, recA | K. Lewis [6] |
| MG1655 ΔrecA | ΔrecA::Kan |  |
| **Plasmids** |  |  |
| pZS*24 | Kan™, lacIq, *E. coli* expression vector | [6] |
| pZS*24tisB | Kan™, lacIq, *tisB* | [6] |
| pCA24N | Cm™, lacIq, *E. coli* expression vector | [6] |
| pCA24NrelA | Cm™, lacIq, *relA* | [27] |

https://doi.org/10.1371/journal.pone.0182122.t001

**Competing interests:** The authors have declared that no competing interests exist.
derivatives, including α-methylcinnamaldehyde, cinnamic alcohol, o-nitrocinamaldehyde, Br-CA, carbicol, phenylpropyl aldehyde, phenylpropanoic acid, and cinnamic acid, were obtained from Tokyo Chemical Co. LTD (TCI). Other reagents were of standard analytical purity.

Persister measurement

*E. coli* MG1655 was cultured in LB overnight and then diluted 1:200 into fresh LB medium. The diluted cultures were incubated at 37˚C with shaking to reach OD600 = 1.0 or stationary growth phase (OD600 = 3.0). The persister formation was evaluated post exposure to 200 μg/ml of Amp and 5 μg/ml of Cipro for 4 hour as described by Orman [25]. Simultaneously, the cells during these phases were treated with CA and its derivatives and the corresponding persisters levels measured. The final concentrations of CA and its derivatives were 5, 10, 50, 100, 200 and 400 μg/ml. The samples were all incubated at 37˚C for 4 hour, and then washed with saline twice. The cell viability was determined using the drop plating method [28]. Spread plating was done to repeat all of the experiments. The assay was repeated at least three times. The hierarchical cluster analysis was performed based on their dose-dependent bactericidal activities using Cluster 3.0 software in the CLUSTER program (http://bonsai.hgc.jp/~mdehoon/software/cluster/software.htm), and the resulting tree figures and cluster color bar were displayed using Java Treeview (http://www.treeview.net/) [29].

Measurement of persisters induced by TisB and RelA overexpression

Overexpression of RelA and TisB in strains MG1655/pCA24NrelA and MG1655/pZS’24tisB, respectively, was used to induce persister formation. Briefly, overnight cultures were inoculated at a ratio of 1:200 into fresh LB. Chloramphenicol (30 μg/ml) and kanamycin (50 μg/ml) were used for maintaining the plasmids pCA24NrelA and pZS’24tisB, respectively. When OD600 value reached 0.4, the culture was supplemented with Isopropyl-β-d-thiogalactoside (IPTG) (0.3 mM for pCA24NrelA and 0.5 mM for pZS’24tisB) to induce relA and tisB overexpression. The cultures were continually incubated until the OD600 value approached 1.0. The cells were collected through centrifugation at 5,000 g for 5 min. The culture medium was discarded and the cells were resuspended in LB medium containing 100 μg/ml of Amp, 5 μg/ml of Cipro and 200 μg/ml of Br-CA respectively, and incubated for 4 hour. Simultaneously, MG1655 containing empty plasmid was included as a control and was treated under the same conditions. Cell viability was determined using the drop plating method [28].

Measurement of persisters induced by chemicals

As described above [10], culture of MG 1655 was grown until OD600~1.0. Then, the cultures were pretreated with Rifä (100 μg/ml for 30 min), Tetra (50 μg/ml for 30 min), or CCCP (20 μg/ml for 3 hour). Following the pretreatment, the cultures were centrifuged to remove the pretreatment compounds and resuspended in fresh LB containing ciprofloxacin (5 μg/ml), ampicillin (100 μg/ml), and Br-CA (200 μg/ml) respectively before being incubated for 4 hour [10]. Cell viability was determined using the drop plating method [28].

MIC and MBC Assays

The minimum inhibitory concentration (MIC) of Br-CA against *E. coli* MG1655 was determined through broth micro dilution assay in a 96-well plate. The bacterial culture during the mid-late exponential phase was prepared and diluted with fresh LB medium until the OD600 value reached 0.05. Subsequently, the diluted culture was transferred into a 96-well plate with
200 μl in each well. Br-CA was added into the wells with final concentrations of 0, 20, 40, 80 and 100 μg/ml. After incubation for 24 hour at 37˚C, OD600 value in each treated well was determined using a microplate reader (Thermo Fischer, United Kingdom). The MIC was defined as the minimum concentration of Br-CA leading to no significant increase of OD600 value. At the same time, the cell viability in each well was assessed using the drop plating method after being washed with saline [28]. The minimum bactericidal concentration (MBC) of Br-CA was defined as the minimum concentration of Br-CA leading to no viable cell determined in the well.

**Hydroxyl-radical scavenging**

According to Kohanski [30] and Piccaro [31], thiourea (Thio) was used as a hydroxyl-radical scavenger and diethylene triaminepenta acetic acid (DTPA) was used as a Fe$^{3+}$ chelating agent to block the hydroxyl-radical production. Thio (final concentration of 100 mM) or DTPA (final concentration of 1 mM) was added into the culture 3 min before the exposure to Br-CA. The effects of Thio and DTPA on the bactericidal activity of Br-CA were assessed in the case of 200 μg/ml of Br-CA against the cells of OD600 = 1.0 or Br-CA at MIC and MBC, respectively against the cells of OD600 = 0.05.

**Data analysis**

All of the experiments in this study were repeated three times, and the results shown are the mean ± SE of the three independent experiments. Significant differences between the treatments were evaluated using SD and one-way analysis of the variance (ANOVA) using SPSS 2.0. The data between the two specific different treatments were compared statistically using Student’s t-test. The differences were considered significant at $P<0.05$.

**Results**

**Bactericidal activities of CA and its derivatives**

The number of persisters increases during the mid-exponential phase [32]. We assessed the bactericidal effects of CA and the eight CA derivatives, i.e. Br-CA, α-methylcinnamaldehyde, cinnamic alcohol, o-nitrocinnamaldehyde, carboxil, phenylpropyl aldehyde, phenylpropanoic acid, and cinnamic acid on *E. coli* MG1655 cells during the exponential phase. As shown in Fig 1, only CA (killing rate: 99.8%), Br-CA (killing rate: 100%), o-nitrocinnamaldehyde (killing rate: 99.8%), and phenylpropyl aldehyde (killing rate: 92.1%) showed significant bactericidal effects at the given concentrations. Through cluster analysis based on their dose-dependent activity, these four compounds were also distributed into the same subgroup, and all of them have aldehyde group (Fig 1). Among these derivatives, Br-CA exhibited the most effective bactericidal activity. When the concentration of Br-CA reached 100 μg/ml, less than 1% of cells survived. Treatment with Br-CA at concentrations over 200 μg/ml completely killed all cells including persisters to common antibiotics.

**MIC and MBC of Br-CA against *E. coli* MG1655**

The MIC value of Br-CA against *E. coli* MG1655 was obtained through a micro-well culture plate assay using serial dilutions of Br-CA. The MBC value was determined by determining the viability of cells in the micro-wells. As shown in Fig 2A and 2B, the MIC value and MBC value of Br-CA against *E. coli* MG1655 were 40 μg/ml and 80 μg/ml, respectively. Br-CA was regarded as bactericidal because the MBC is no more than four times of the MIC according to French [33]. Based on the MIC and MBC of Br-CA, 200 μg/ml of Br-CA (concentration of...
5xMIC) was chosen to investigate its potential to eradicate the persister cells in *E. coli* MG1655.

**Br-CA eradicated persisters in stationary phase**

It has been well accepted that more persisters are formed during the stationary phase [34], which was supported by our results that in stationary phase treated by Amp the level of persisters increased to approximately 1% of the population, while it remained at low level when cells were treated by Cipro (Fig 3). Br-CA displayed a different dose-dependent activity against stationary phase *E. coli* cells as compared to exponential phase cells, but it was able to eradicate the stationary phase persisters when its concentration reached 400 μg/ml.

**Br-CA eradicated persisters induced by toxin overexpression**

Since ppGpp can induce the formation of persisters [9], we employed the RelA overexpression strain MG1655/pCA24NrelA in which ppGpp was over produced to induce persister formation. Our results showed that overproduction of ppGpp induced more than 430-fold and 1890-fold of persisters induced to Amp and Cipro, respectively (Fig 4A). Br-CA completely eradicated all of the cells (killing rate: 100%) including persisters induced by ppGpp. The toxin TisB in SOS can also induce persister formation through the elimination of cellular ATP [6]. In the present study, the MG1655/pZS₂4tisB strain in which TisB was overexpressed led to 4.9-fold and 17-fold increases in persister formation as compared to those induced by Amp and Cipro, respectively (Fig 4B). Similar to its efficiency against the strain pCA24NrelA, Br-
CA also eradicated all pZS'24 \( tisB \) cells (killing rate: 100%). Surprisingly, a small fraction of cells containing empty vectors survived Br-CA treatment in the presence of IPTG (Fig 4A and 4B), indicating that IPTG probably helped cell survival after exposure to Br-CA. Indeed, in absence of IPTG, persisters were eradicated by Br-CA while only slight changes occurred in persisters levels obtained after treatment by Amp and Cipro. In addition, we also determined...
the effects of IPTG on persister formation in wild type strains. In comparison to the control group (without IPTG), addition of IPTG did not significantly change the levels of persisters when treated by Amp and Cipro, whereas it substantially increased persisters levels when treated by Br-CA (Fig 4C).

Potential of Br-CA against chemical-induced persister formation

Rifampin (halting transcription), tetracycline (halting translation), and CCCP (halting ATP synthesis) have been used to mimic type II and TisB-like toxins to induce persister formation, respectively [10]. In our current study, these three compounds all led to more than 1000-fold increase in the persister levels when treated by Amp or Cipro (Fig 5). Br-CA completely eradicated all cells (killing rate, 100%) under induction by CCCP (Fig 5), which was in line with the results from the overexpression of TisB (Fig 4B). However, Br-CA failed to eradicate the cells under the induction of Rifa and Tetra (Fig 5), and it was obvious that persisters induced by Rifa or Tetra were also tolerant to Br-CA (Fig 5). This is inconsistent with the results from the overproduction of ppGpp, indicating that the mechanism underlying persister formation by Rifa and Tetra was different from the ppGpp pathway. In strain ΔrecA (a mutant with the deletion of recA to block SOS), persists levels after treatment by Br-CA and pre-treatment by Rifa and Tetra decreased by 790-fold and 1906-fold, respectively, as compared to the wild type (Fig 5).

The role of ROS in the bactericidal activity of Br-CA

Reactive oxygen species (ROS) are known to cause important damage to the cell, which can lead to cell death [35, 36]. In order to investigate whether ROS was involved in the high bactericidal activity of Br-CA, thiourea (hydroxyl-radical scavenger) and the iron chelator DTPA (blocking hydroxyl-radical production) were employed. As shown in Fig 6, the addition of Thio or DTPA either did not affect the eradication of exponential cells of E. coli under Br-CA treatment. Further, the effects of Thio or DTPA on the bactericidal activity of Br-CA at MBC (80 μg/ml) and MIC (40 μg/ml) was also investigated, respectively. The results showed that Thio or DTPA did not antagonize the bactericidal effects of Br-CA at MBC (80 μg/ml), and they even promoted bacterial death by Br-CA at MIC (40 μg/ml) (Fig 6). These results suggested that ROS could play a protective role rather than causing cell death in E. coli after exposure to Br-CA.
As a derivative of the essential oil CA, Br-CA is capable of causing slightly non-specific inflammatory irritating effects on the skin and eye [37]. According to the safety data sheet of the products by Sigma-Aldrich Co., Br-CA is defined as chemical of the 4th grade of acute toxicity in oral administration (LD50 > 2000 mg/Kg). Br-CA has been reported to possess a dose-independent genotoxic and mutagenic potential in bacteria [38], but to date, no DNA-adducts of Br-CA or any clear carcinogenic activities have been found. The present study demonstrated Br-CA had the most powerful bactericidal activity against *E. coli* among CA and its derivatives, with especially high efficiency in the eradication of persisters. Balaban and colleagues [39] have defined two types of *E. coli* persisters based on their growth-rate and formation period, i.e. type I persisters forming from the stationary phase and type II persisters that are continuously generated during growth. Type I persisters especially reach 1% of the entire population [34]. It was inspiring that Br-CA could eradicate both type I and type II persisters. Recent studies have indicated that the anticancer drugs mitomycin C and cisplatin eradicate persisters by cross-linking their DNA [14, 40]. However, this kind of bactericidal mechanism also leads to significant side effects [41, 42], and they are also very costly. In comparison to them, Br-CA has a low price, showing potential to be applied in the treatment of infection. However, there is very
scarce data regarding Br-CA toxicity, and we should be cautious in its application in the treatment of infections.

The ppGpp has been recognized as a central regulator that inhibits translation and cell growth, thereby inducing persistence through activation of toxins [9]. Our results showed that Br-CA completely eradicated the persisters induced by the overproduction of ppGpp, but not that induced by Rif or Tetra. Since the antimicrobial roles of Rif and Tetra have been targeted at the RNA polymerase and 30S ribosomal subunit, respectively [43, 44], we proposed that ppGpp-induced persister formation is independent of the inhibition of transcription or translation. The recent study has shown that ppGpp couples transcription to DNA repair in *E. coli*, which contributes to its survival under genotoxic stress [45]. Therefore, SOS may be involved in ppGpp-induced persister formation to some antibiotics. Blocking SOS by deleting recA significantly decreased Rif- or Tetra-induced persisters to Br-CA as compared to the wild-type (Fig 5), which suggests the involvement of SOS in the induction of persister formation by Rif or Tetra. Dörr et al., [6] have identified a SOS-associated persister formation pathway through the activation of

**Fig 6. Thiourea and DTPA have no effect on the eradication of persisters by Br-CA.** The initial bacterial suspension (OD600 = 0.05 or OD600 = 1.0) was pretreated with or without Thio or DTPA iron chelator and then exposed to different concentrations Br-CA. Cell survival percentage after treatment was calculated based on the colony counts on the LB agar plates. The arrows indicated that cells were killed completely by Br-CA and different colors showed different treatments. Each bar was indicated as the means of three replicates ± standard deviation. Survival post treatment with Br-CA alone at the given concentration were set as the control in the corresponding group. Asterisks indicate a significant difference from their controls (*P*<0.05).

https://doi.org/10.1371/journal.pone.0182122.g006
TisB, leading to ATP depletion. However, Br-CA effectively eradicated the persisters induced by either TisB overexpression or CCCP. This suggested that there might be other elements in SOS in addition to TisB that induce the persister formation. IPTG increased the cells that survived the Br-CA treatment, whereas the overexpression of toxins eliminated this protective effect of IPTG, indicating that toxins overexpression leading to persister formation to Amp and Cipro could be helpful in the eradication of persisters by Br-CA. Taken together, we propose that the mechanisms underlying persister formation should be more diverse than we thought previously, and Br-CA might be considered for the application in the infection treatment to overcome the persister problem if its toxicity could be proved to be acceptable in the future.

CA activates TRPA1 by covalently binding via Michael addition, leading to a Ca\(^{2+}\) influx [46]. Our previous study obtained similar results in which the stimulation of a transient Ca\(^{2+}\) efflux was involved in the CA-induced growth inhibition of \textit{P. capsici} [20]. These studies imply that the potential of \(\alpha, \beta\)-unsaturated bond undergoing Michael addition is directly associated with its antifungal activity. Br-CA has a very active \(\alpha, \beta\)-unsaturated bond to allow for Michael addition [38], which seemed to correspond to its highest bactericidal activity. Exceptionally, \(\alpha\)-methylcinnamaldehyde with an \(\alpha, \beta\)-unsaturated bond did not display any bactericidal activity at any given concentrations, whereas phenylpropyl aldehyde without an \(\alpha, \beta\)-unsaturated bond exerted bactericidal activity. This indicated that the bactericidal activity of CA derivatives was probably independent of Michael addition. In our present study, neither Thio (hydroxyl-radical scavenger) nor DPTA (Fe\(^{3+}\) chelator to block the hydroxyl-radical) affected the bactericidal efficiency of Br-CA against \textit{E. coli}, which suggested a novel bactericidal mechanism of Br-CA independent of ROS. Taken together, our results indicated that Br-CA had efficient bactericidal activity with mechanisms different from common antibiotics, which implemented plant resources (e.g. CA and Br-CA) to be developed to tackle the antibiotics resistance crisis.

**Acknowledgments**

The authors are grateful to Professor Kim Lewis at Northeastern University for sharing the strains of MG1655, MG1655\(\Delta\text{recA}\), MG1655/pZS\(^{24}\), and MG1655/pZS\(^{24}\)\(\text{tisB}\), as well as Professor Thomas K. Wood at Pennsylvania State University for plasmids pCA24N and pCA24N-\(\text{relA}\). This work was supported by funding from the Program for Science and Technology Innovation Team in Universities of Henan Province (16IRTSTHN007), Key Scientific and Technological Projects of Henan Province (NO. 152102110086) and Project of Youth Backbone Teachers of Colleges and Universities in Henan Province (2014GGJS-101).

**Author Contributions**

**Data curation:** Liangbin Hu.

**Formal analysis:** Wei Zhou, Yonghua Qi, Hongmei Ning, Jian Chen, Haizhen Mo.

**Investigation:** Qingshan Shen, Wei Zhou, Jian Chen.

**Methodology:** Qingshan Shen, Liangbin Hu.

**Project administration:** Liangbin Hu, Yonghua Qi, Hongmei Ning, Haizhen Mo.

**Supervision:** Liangbin Hu, Haizhen Mo.

**Validation:** Liangbin Hu, Haizhen Mo.

**Writing – original draft:** Qingshan Shen, Liangbin Hu.

**Writing – review & editing:** Liangbin Hu, Yonghua Qi, Hongmei Ning, Haizhen Mo.
References

1. Ayrapetya N, Williams TC, Oliver JD. Bridging the gap between viable but non-culturable and antibiotic persistent bacteria. Trends in microbiology. 2015; 23(1):7–13. https://doi.org/10.1016/j.tim.2014.08.004 PMID: 25449050
2. Bigger J. Treatment of staphylococcal infections with penicillin by intermittent sterilisation. The Lancet. 1944; 244(6320):497–500.
3. Wiuff C, Zappala R, Regoes R, Garner K, Baquero F, Levin B. Phenotypic tolerance: antibiotic enrichment of noninherited resistance in bacterial populations. Antimicrobial agents and chemotherapy. 2005; 49(4):1483–94. https://doi.org/10.1128/AAC.49.4.1483-1494.2005 PMID: 15793130
4. Cohen NR, Lobritz MA, Collins JJ. Microbial Persistence and the Road to Drug Resistance. Cell Host & Microbe. 2013; 13(6):632–42.
5. Bernier SP, Lebeaux D, Defrancesco AS, Valomon A, Soubigou G, Coppée JY, et al. Starvation, together with the SOS response, mediates high biofilm-specific tolerance to the fluoroquinolone ofloxacin. Plos Genetics. 2013; 9(1):e1003144. https://doi.org/10.1371/journal.pgen.1003144 PMID: 23300476
6. Dörr T, Vulic M, Lewis K. Ciprofloxacin causes persister formation by inducing the TisB toxin in Escherichia coli. Public Library of Science; 2010. 1000317 p.
7. Vega NM, Allison KR, Khalil AS, Collins JJ. Signaling-mediated bacterial persister formation. Nature Chemical Biology. 2012; 8(9):431. https://doi.org/10.1038/nchembio.915 PMID: 22426114
8. Wu Y, Vulic M, Keren I, Lewis K. Role of Oxidative Stress in Persister Tolerance. Antimicrobial Agents & Chemotherapy. 2012; 56(9):4922.
9. Maisonneuve E, Gerdes K. Molecular Mechanisms Underlying Bacterial Persisters. Cell. 2014; 157(3):539. https://doi.org/10.1016/j.cell.2014.02.050 PMID: 24766804
10. Kwan BW, Valenta JA, Benedik MJ, Wood TK. Arrested protein synthesis increases persister-like cell formation. Antimicrobial agents and chemotherapy. 2013; 57(3):1468–73. https://doi.org/10.1128/AAC.02135-12 PMID: 23295927
11. Chen X, Zhang M, Zhou C, Kallenbach NR, Ren D. Control of bacterial persister cells by Trp/Arg-containing antimicrobial peptides. Appl Environ Microbiol. 2011; 77(14):4878–85. https://doi.org/10.1128/AEM.02440-10 PMID: 21622798
12. Conlon BP, Nakayasu ES, Fleck LE, LaFleur MD, Isabella VM, Coleman K, et al. Activated ClpP kills persisters and eradicates a chronic biofilm infection. Nature. 2013; 503(7476):365–70. https://doi.org/10.1038/nature12790 PMID: 24226776
13. Marques CN, Morozov A, Planzos P, Zelaya HM. The fatty acid signaling molecule cis-2-decenoic acid increases metabolic activity and reverts persister cells to an antimicrobial-susceptible state. Appl Environ Microbiol. 2014; 80(22):6976–91. https://doi.org/10.1128/AEM.01576-14 PMID: 25192989
14. Kwan BW, Chowdhury N, Wood TK. Combating bacterial infections by killing persister cells with mitomycin C. Environ Microbiol. 2015; 17(11):4406–14. https://doi.org/10.1111/1462-2920.12873 PMID: 25858802
15. Moreira W, Aziz DB, Dick T. Boromycin Kills Mycobacterial Persisters without Detectable Resistance. 2016. https://doi.org/10.3389/fmicb.2016.00199 PMID: 26941723
16. Pan J. Controlling Bacterial Persister Cells and Biofilms by Synthetic Brominated Furanones. Dissertation, Syracuse University. 2013.
17. Zhang Y. Persisters, persistent infections and the Yin—Yang model. Emerging Microbes & Infections. 2014; 3(1):e3.
18. Teng L, Zu Q, Li G, Yu T, Job KM, Yang X, et al. Herbal medicines: challenges in the modern world. Part 3. China and Japan. Expert review of clinical pharmacology. 2016; 1:1–9. https://doi.org/10.1080/17512433.2016.1195263 PMID: 27232545
19. Upadhyay A, Venkitanarayanan K. In vivo efficacy of trans-cinnamaldehyde, carvacrol, and thymol in attenuating Listeria monocytogenes infection in a Galleria mellonella model. Journal of Natural Medicines. 2016; 70(3):667–72. https://doi.org/10.1007/s11418-016-0990-4 PMID: 27094514
20. Hu L, Wang D, Liu L, Chen J, Xue Y, Shi Z. Ca²⁺ efflux is involved in cinnamaldehyde-induced growth inhibition of Phytophthora capsici. PLoS one. 2013; 8(10):e76264. https://doi.org/10.1371/journal.pone.0076264 PMID: 24098458
21. Budri P, Silva N, Bonzagli E, Fernandes A, Araújo J, Doyama J, et al. Effect of essential oils of Syzygium aromaticum and Cinnamomum zeylanicum and their major components on biofilm production in Staphylococcus aureus strains isolated from milk of cows with mastitis. Journal of dairy science. 2015; 98(9):5899–904. https://doi.org/10.3168/jds.2015-9442 PMID: 26142866
22. Shafreen B, Mohmed R, Selvaraj C, Singh SK, Karutha Pandian S. In silico and in vitro studies of cinnamaldehyde and their derivatives against LuxS in Streptococcus pyogenes: effects on biofilm and virulence genes. Journal of Molecular Recognition. 2014; 27(2):106–16. https://doi.org/10.1002/jmr.2339 PMID: 24436128

23. Nostro A, Scalfaro R, Botta L, Filocamo A, Marino A, Bisignano G. Effect of temperature on the release of carvacrol and cinnamaldehyde incorporated into polymeric systems to control growth and biofilms of Escherichia coli and Staphylococcus aureus. Biofouling. 2015; 31(8):639–49. https://doi.org/10.1080/08927014.2015.1079703 PMID: 26362127

24. Römling U, Kjelleberg S, Normark S, Nyman L, Uhlin BE, Åkerlund B. Microbial biofilm formation: a need to act. Journal of Internal Medicine. 2014; 276(2):98–110. https://doi.org/10.1111/joim.12242 PMID: 24796496

25. Orman MA, Brynildsen MP. Persister formation in Escherichia coli can be inhibited by treatment with nitric oxide. Free Radical Biology and Medicine. 2016; 93:145–54. https://doi.org/10.1016/j.freeradbiomed.2016.02.003 PMID: 26849946

26. Sambrook J, Fritsch EF, Maniatis T. Molecular cloning: a laboratory manual. New York, Cold Spring Harbour Press. 1990.

27. Kitagawa M, Ara T, Arifuzzaman M, Iokanakamichi T, Inamoto E, Toyonaga H, et al. Complete set of ORF clones of Escherichia coli ASKA library (A Complete Set of E. coli K-12 ORF Archive): Unique Resources for Biological Research. DNA Research. 2005; 12(5):291. https://doi.org/10.1093/dnares/dsi012 PMID: 16769691

28. Herigstad B, Hamilton MA, Heersink J. How to optimize the drop plate method for enumerating bacteria. Journal of Microbiological Methods. 2001; 44(2):121–9. PMID: 11165341

29. de Hoon MJ, Imoto S, Nolan J, Miyano S. Open source clustering software. Bioinformatics (Oxford, England). 2004; 20(9):1453–4. https://doi.org/10.1093/bioinformatics/bth078 PMID: 14871861

30. Kohanski MA, Dwyer DJ, Hayete B, Lawrence CA, Collins JJ. A common mechanism of cellular death induced by bactericidal antibiotics. Cell. 2007; 130(5):797–810. https://doi.org/10.1016/j.cell.2007.06.049 PMID: 17803904

31. Piccaro G, Pietrafitta D, Giannoni F, Mustazzolo A, Fattorini L. Rifampin Induces Hydroxyl Radical Formation in Mycobacterium tuberculosis. Antimicrobial Agents and Chemotherapy. 2014; 58(12):7527–33. https://doi.org/10.1128/AAC.03169-14 PMID: 25288092

32. Keren I, Kaldalu N, Spoering AL, Wang Y, Lewis K. Persister cells and tolerance to antimicrobials. FEMS Microbiology Letters. 2004; 230(1):13–8. https://doi.org/10.1016/s0378-1097(03)00856-5 PMID: 14734160

33. French GL. Bactericidal agents in the treatment of MRSA infections—the potential role of daptomycin. The Journal of antimicrobial chemotherapy. 2006; 58(6):1107–17. https://doi.org/10.1093/jac/dkl393 PMID: 17040922

34. Lewis K. Persister cells, dormancy and infectious disease. Nature reviews Microbiology. 2007; 5(1):48–56. https://doi.org/10.1038/nrmicro1557 PMID: 17143318

35. Albesa I, Becerra MC, Battán PC, Páez PL. Oxidative stress involved in the antibacterial action of different antibiotics. Biochemical and biophysical research communications. 2004; 317(2):605–9. https://doi.org/10.1016/j.bbrc.2004.03.085 PMID: 15063800

36. Schenk B, Fulda S. Reactive oxygen species regulate Smac mimetic|[[sol]]|TNF|[[alpha]]| induced necrototic signaling and cell death. Oncogene. 2015; 34(47):5796. https://doi.org/10.1038/onc.2015.35 PMID: 25867066

37. Lu DBJ, He Q, Wang J, Chen Y, Yin M. Studies on the toxicities of alpha-bromocinnamaldehyde and structure-effect relationship. Carcinogenesis Teratogenesis & Mutagenesis. 1992; 4:22–5.

38. Eder E, Deiningger C, Deiningger D, Weinfurtner E. Genotoxicity of 2-halocinnamaldehydes in bacterial assays. Induction of SOS repair and frame-shift mutation. Mutagenesis. 1994; 9(5):473–6. PMID: 7837982

39. Balaban NQ, Merrin J, Chait R, Kowalik L, Leibler S. Bacterial Persistence as a Phenotypic Switch. Science. 2004; 305(5690):1622–5. https://doi.org/10.1126/science.1099390 PMID: 15308767

40. Chowdhury N, Wood TL, Martinezvazquez M, Garciacontreras R, Wood TK. DNA-crosslinker cisplatin eradicates bacterial persister cells. Biotechnology and Bioengineering. 2016; 113(9):1984–92. https://doi.org/10.1002/bit.25963 PMID: 26914280

41. Cicek D, Cobanoglu B, Inci R, Dertioglu SB, Kokcam I, Elirkian T. A very rare side effect of mitomycin-C: Psoriasiform drug eruption. International Journal of Dermatology. 2013; 52(12):1572–4. https://doi.org/10.1111/j.1365-4632.2011.05306.x PMID: 24261729

42. Levi JA, Aroney RS, Dalley DN. Haemolytic anaemia after cisplatin treatment. British Medical Journal. 1981; 282(6281):2003–4. PMID: 6788166
43. Brodersen DE, Clemons WM Jr., Carter AP, Morgan-Warren RJ, Wimberly BT, Ramakrishnan V. The structural basis for the action of the antibiotics tetracycline, pactamycin, and hygromycin B on the 30S ribosomal subunit. Cell. 2000; 103(7):1143–54. PMID: 11163189

44. Floss HG, Yu TW. Rifamycin-mode of action, resistance, and biosynthesis. Chemical reviews. 2005; 105(2):621–32. https://doi.org/10.1021/cr030112j PMID: 15700959

45. Kamarthapu V, Epshtein V, Benjamin B, Proshkin S, Mironov A, Cashel M, et al. ppGpp couples transcription to DNA repair in E. coli. Science. 2016; 352(6288):993–6. https://doi.org/10.1126/science.aad6945 PMID: 27199428

46. Macpherson LJ, Dubin AE, Evans MJ, Marr F, Schultz PG, Cravatt BF, et al. Noxious compounds activate TRPA1 ion channels through covalent modification of cysteines. Nature. 2007; 445(7127):541–5. https://doi.org/10.1038/nature05544 PMID: 17237762