



## Original Research Article

# Empowering Therapeutic Strategies against Methicillin-Resistant *Staphylococcus Aureus* Riboswitch: Unveiling the Potential of Small Molecules and Antisense Oligonucleotides through *In Silico* Analysis

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## ABSTRACT

Methicillin-resistant *Staphylococcus aureus* (MRSA) poses an increasingly alarming threat to global public health, characterized by its escalating antibiotic resistance and heightened virulence. This dire situation necessitates the exploration of novel and hitherto uncharted targets for effective pathogen control. The primary objective of our current investigation is to identify the most promising riboswitch as a druggable target and assess its potential inhibition through both small molecules and antisense oligonucleotides (ASO) *in silico*. I conducted an exhaustive search for the most plausible druggable riboswitch using the RiboScan tool, subsequently subjecting its sequence to rigorous docking studies with ten distinct resveratrol derivatives through the PatchDock algorithm. In addition, we harnessed the power of ASO by designing and hybridizing five ASO sequences to silence the identified riboswitch target effectively. This comprehensive analysis pinpointed the glmS riboswitch as the most promising candidate for targeted intervention. Molecular docking results uncovered the exceptional inhibitory potential of resveratrol derivatives, with particular emphasis on ZINC000100827960, which emerged as a lead natural inhibitor. Furthermore, the ASO sequences, meticulously designed for their intended purpose, exhibited compelling efficacy in terms of hybrid free energy, RNA-RNA docking interactions, and a remarkably low probability of causing off-target effects within the human transcriptome. Taken together, our findings underscore the significant promise of both resveratrol derivatives and the ASO candidates as potent inhibitors of the glmS riboswitch in MRSA. These results warrant not only further consideration, but also rigorous experimental validation to pave the way for innovative strategies in combating the menace posed by MRSA.

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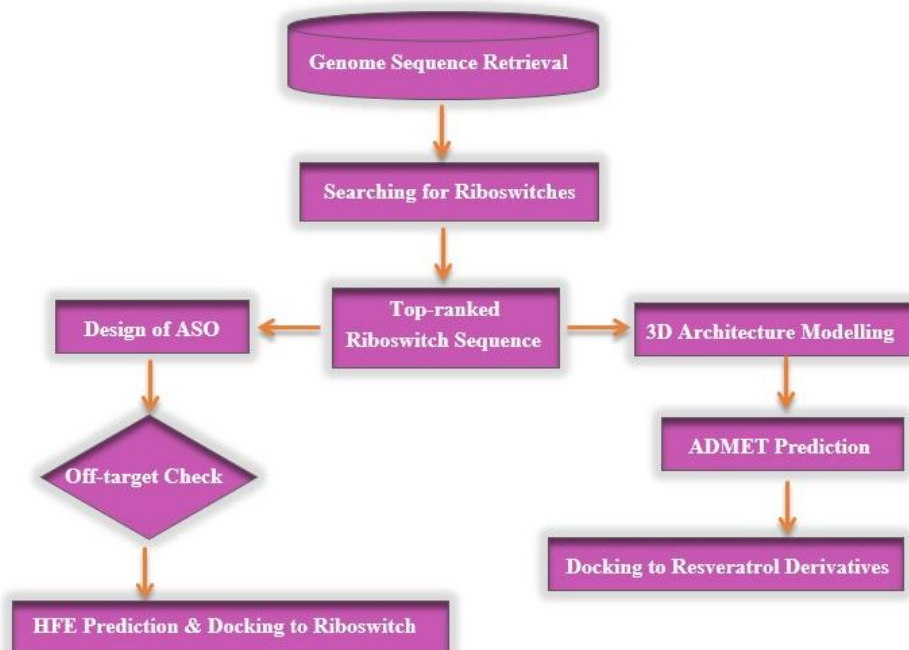
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**GRAPHICAL ABSTRACT**


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## Introduction

*Staphylococcus aureus* is a pathogenic bacterium that results in many organ dysfunctions, including superficial (skin) and deep ones (joints and cardiac tissues). This anaerobic, Gram-positive bacterium causes both community-acquired as well as hospital-acquired serious infections [1]. Due to the misuse of antibiotics against such highly virulent pathogen, the medical community encounters the emergence of antibiotic-resistant strains, even to the highly potent methicillin antibiotic (methicillin-resistant *S. aureus*, MRSA). Nowadays, more than 100 MRSA strains are classified as major of concern in USA. This led the World Health Organization (WHO) to deem MRSA as an endemic due to limited available therapeutics [2]. Vancomycin was an emergency-based therapeutic option to deal with MRSA strains but, again, vancomycin-resistant *S. aureus* (VRSA) was also developed and now is widely distributed in USA, India, Pakistan, and Iran [3]. It is estimated that approximately 5% of

population carry MRSA. MRSA and VRSA show notorious virulence and lead to life-threatening consequences [4]. Accordingly, the search for new generation of novel targets for controlling MRSA is significantly demanding.

Bacterial genomes are classified into two main components: the core genome and the accessory genome. The core genome is composed of genes that are present in all isolates and contains essential genetic information related to cellular metabolism and replication. It makes up approximately 75% of the 2.8 Mb genome of *S. aureus* and is highly conserved among strains [5]. On the other hand, the accessory genome, which comprises about 25% of the total *S. aureus* genome, is more variable and often more strain-specific than the core genome. It consists of mobile genetic elements such as pathogenicity islands, bacteriophages, chromosomal cassettes, transposons, and plasmids acquired by horizontal transfer between strains. This is where mediators of virulence, immune evasion, and antibiotic resistance are commonly found [6,7].

One of the less-studied targets is the bacterial riboswitch. Riboswitch is a structured RNA transcribed from the 5'-untranslated region (5'-UTR) of a gene and controls gene expression from which synthesized. In other words, riboswitch is a mediator that senses the changes of some essential amino acids and metabolites in response to environmental fluctuations to increase/decrease the expression of metabolite-synthesizing gene [8].

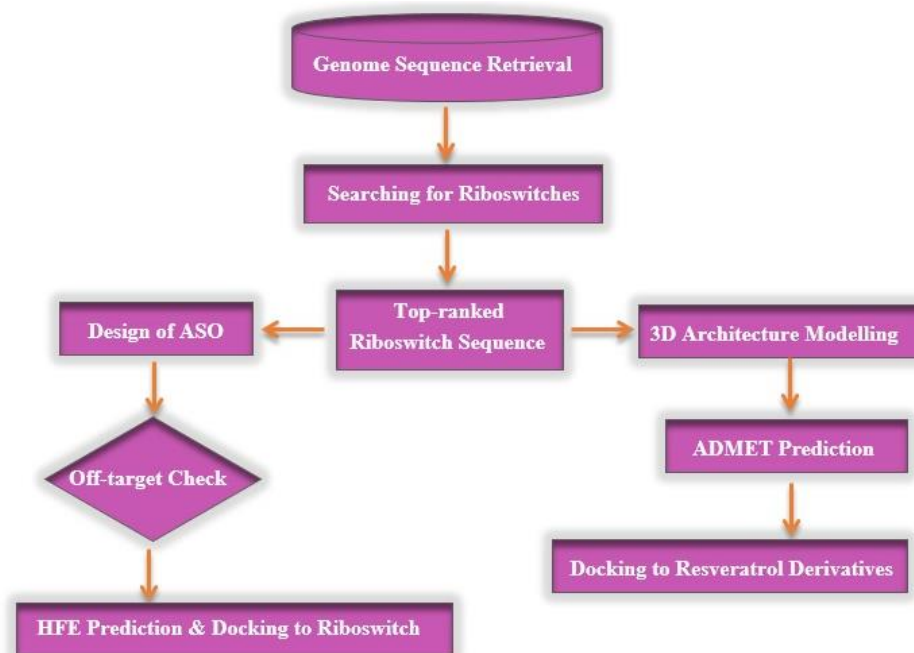
Riboswitch offers many merits as an antibacterial target including (i) function independently of proteins or enzymes, (ii) exclusively present in prokaryotes, (iii) relatively small size (often < 100 nucleotides), and (iv) located in the 5'-UTR which conceives less impact on the coding region of the related genes [9]. With respect to structure-to-function relationship, riboswitch is composed of 2 domains: Ligand-binding domain (aptamer) and regulatory domain. Once the ligand got bound to the corresponding riboswitch in the aptamer domain, it induces conformational modifications to the overall structure facilitating, thus it is binding to the cognate gene or mRNA. This in

turn blocks or triggers transcription or translation of the related gene (ON/OFF-switching) [10]. PKZ18 was demonstrated to target T-box riboswitch of MRSA with no substantial cytotoxicity toward eukaryotic cells after exposure for 24 h, but its analogues improved the specificity minimizing the cytotoxicity [11]. Similarly, with an MIC (minimum inhibitory concentration) = 0.5 mg/mL, Ribocil-C was able to diminish the riboflavin uptake and its binding to flavin mononucleotide (FMN) riboswitch. This confirmed the antibacterial action versus MRSA [12].

The computational analysis of small molecules docking and antisense oligonucleotide (ASO) versus MRSA riboswitches is scarce. Hence, the present study purposes to inhibit MRSA riboswitch by small molecules and ASO.

## Materials and methods

The outlined methodology used in the current theoretical study is shown in Figure 1. However, the details of every step are elucidated in detail below.



**Figure 1.** Flowchart of the methodology employed in the present study.

### Retrieval of sequence

The MRSA sequence was downloaded from the national center for biotechnology information (NCBI) [13]. The complete genome of MRSA strain M92 has an accession ID: CP015447.2 which contains ~ 3 million bp in total.

### Searching for riboswitches

As an input, the full genome FASTA sequence of MRSA was uploaded to RiboScan tool (<http://service.iiserkol.ac.in/~riboscan/index.html>) [14] in so as to get the possible riboswitches. The tool besides finding the possible riboswitches, also ranks them by a score and detect whether the found riboswitch in the coding or complementary strands.

The tool uses a method based on profile Hidden Markov Models (pHMMs) for detecting riboswitches.

### Prediction of secondary and tertiary architectures

The RiboScan tool provides also the overall secondary structure of the obtained riboswitch classes. In addition, the tertiary structure of the top-rank riboswitch was predicted from the provided sequence using 3dRNA webserver (<http://biophy.hust.edu.cn/new/3dRNA/>) [15]. The chosen procedure was "optimize" in order to get the best generated model. The dot-bracket pattern of the provided sequence was obtained from RNAfold (<http://rna.tbi.univie.ac.at/cgi-bin/RNAWebSuite/RNAfold.cgi>) [16].

### Molecular docking

The obtained 3D structure from 3dRNA webserver was used further for molecular docking by PatchDock platform (<https://bioinfo3d.cs.tau.ac.il/PatchDock/>) [17]. The riboswitch structure was prepared by adding polar hydrogen and assigning charge prior to docking. This receptor was docked against 10 resveratrol derivatives in which the previous reports confirmed their potency against

MRSA and its biofilm [18-20] and compared with the reference substrate, glucosamine-6-phosphate. The SDF files of the ligands were downloaded from ZINC20 dataset [21]. Only the best model, along with the native substrate, will be selected for investigation of RNA-ligands interactions by BINANA 2.0 server (<https://durrantlab.pitt.edu/binana/>) [22].

### ADMET profiling

The absorption, distribution, metabolism, excretion, and toxicity (ADMET) of the best ligand will be predicted via admetSAR 2 web portal (<http://lmm.d.ecust.edu.cn/admetSAR2>) [23]. The server predicts Lipinski's rule of five (Ro5) as well as other toxicity parameters via inputting the SMILES of the ligand.

### Design of antisense oligonucleotides (ASO)

PFRED [24] was utilized for the design of the potential ASO versus the riboswitch sequence. The program provided 199 ASOs which filtered using 4 criteria: (i) avoided 5'-untranslated region (UTR), (ii) avoided 3'-UTR, (iii) avoided polyA, and (iv) predicted ASO efficacy score > 10. The best ASO candidates will be tested for off-target with human genome/transcriptome by BLAST database (<https://www.ncbi.nlm.nih.gov/geo/query/blast.html>) [25]

### Prediction of hybrid free energy

To evaluate the lowest hybrid free energy (HFE) upon folding the obtained ASO and their cognate strand, DuplexFold tool [26] (<https://rna.urmc.rochester.edu/RNAstructureWeb/Servers/DuplexFold/DuplexFold.html>) was used. However, the results were validated via RNA-RNA molecular docking tool HNADOCK (<http://huanglab.phys.hust.edu.cn/hnadock/>) [27].

## Results

### Searching for riboswitches

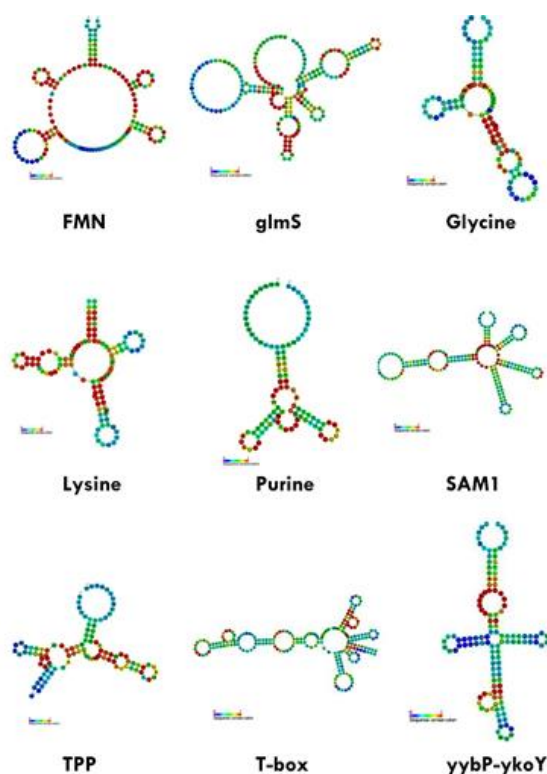
RiboScan output listed 9 riboswitches in the main strand and the complementary strand with a score range from 11.9 to 100.1 (Table 1). The great majority of riboswitches were found to belong to T-box class (10 riboswitches) followed by SAM1 class (3 riboswitches), Lysine, FMN, and

Lysine (2 riboswitches each) whereas the remaining (glmS, Glycine, purine, and yybP-ykoY) had only one riboswitch. However, glmS exhibited the highest score (100.1) with an E-value of  $6.4e-33$  making it the most prominent riboswitch to be targeted in MRSA. Accordingly, only glmS riboswitch (GR) will be considered for further evaluation. The secondary architecture of the obtained all classes is illustrated in Figure 2.

**Table 1.** The obtained riboswitches from the provided MRSA genome sequence

Riboswitch family	Strand	Score (%)	E-value	Region from	Region to	
1	FMN	CS	61.3	5.9e-21	1573870	1574004
		CS	65.4	3e-22	1877643	1877778
2	glmS	CS	100.1	6.4e-33	2420135	2420351
3	Glycine	CS	12.0	8.7e-06	1625117	1625210
4	Lysine	MS	12.8	5.5e-06	1448360	1448525
		CS	57.9	7.5e-20	1761320	1761495
5	Purine	MS	46.4	1.6e-16	437274	437375
6	SAM1	MS	37.8	1.2e-13	15939	16037
		MS	45.5	4.4e-16	908927	909030
		CS	45.9	2.8e-16	1894421	1894531
		CS	44.2	9.5e-16	2592584	2592679
7	TPP	CS	25.7	4.6e-10	1104121	1104220
		CS	34.9	6e-13	2363329	2363429
8	T-box	MS	49.9	3.1e-17	12478	12693
		MS	11.9	1.4e-05	1161904	1162227
		MS	23.9	3e-09	1217395	1217569
		MS	22.6	7.8e-09	1421786	1421962
		MS	19.5	6.8e-08	1421990	1422176
		CS	59.6	3.4e-20	415564	415933
		CS	40.1	3.4e-14	1697153	1697335
		CS	15.3	1.4e-06	1711620	1711872
		CS	71.8	6.3e-24	1744195	1744495
		CS	25.3	1.2e-09	1763699	1763905
9	yybP-ykoY	MS	28.2	9.8e-11	1042326	1042465

\*CS: Complementary strand (3'-5'), MS: Main strand (5'-3)



**Figure 2.** The provided secondary structure of the main riboswitch families found in the genome of MRSA.

### Molecular docking

Ten resveratrol derivatives were selected as potential inhibitors to GR. PatchDock output revealed that all of the tested ligands exhibited higher binding score and atomic contact energy than the native substrate glucosamine-6-phosphate, the best of which was

ZINC000100823225 with a docking score 6326 and atomic contact energy -359.20 kcal/mol (Table 2).

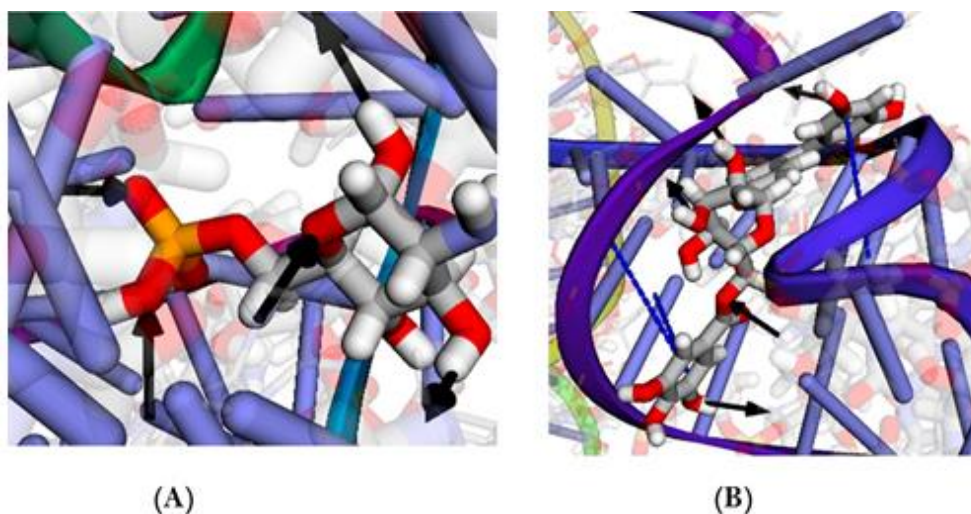
BINANA 2.0 sever unveiled the formation of 5 H-bonds with this ligand as well as 2  $\pi$ - $\pi$  stacking interactions. Similarly, glucosamine-6-phosphate had only 5 H-bonds (Figure 3).

**Table 2.** PatchDock findings of resveratrol derivatives and the cogante ligand against GR

Ligand	PatchDock		
	Score (kcal/mol)	Area (Å)	ACE (kcal/mol)
ZINC000035653092	4566	511.60	-357.69
ZINC000040977346	5120	618.00	-409.09
ZINC000015112534	5270	633.20	-444.92
ZINC000100827965	6232	717.20	-454.77
ZINC6787	4148	452.60	-289.77
ZINC000095620822	4116	459.50	-329.16
ZINC000100827960	6132	735.40	-442.84
ZINC000085612047	6046	725.20	-521.44
ZINC000100823225	6326	717.40	-359.20
ZINC000003978779	4632	516.10	-336.38
Glucosamine-6-phosphate	3328	377.60	-164.63

\*ACE: Atomic contact energy





**Figure 3.** Cognate ligand (A) and ZINC000100823225 and (B) interaction with GR as unveiled by BINANA 2.0 server. Solid black arrow indicates hydrogen bond from donor to acceptor while dashed blue line indicates  $\pi$ - $\pi$  stacking.

**Table 3.** ADMET profile of the best ligand as predicted by admetSAR 2

	Attribute	Result
Ro5	Molecular Weight	542.49
	AlogP	1.43
	H-Bond Acceptor	12
	H-Bond Donor	8
	Rotatable Bonds	7
Toxicity	Carcinogenicity	Negative
	Eye corrosion	Negative
	Eye irritation	Negative
	Hepatotoxicity	Negative
	Nephrotoxicity	Negative
	Respiratory toxicity	Negative
	Skin irritation	Negative
	skin sensitisation	Negative

#### ADMET profile

It is prominent that the best docked ligand (ZINC000100823225) was predicted to pose a good pharmacokinetic profile albeit it displayed a violation to Ro5 in terms of increased H-bond acceptor atoms (12) which was higher than the restricted level (< 10). Otherwise, the ligand is a lead candidate. In addition, the ligand also showed neither potential toxicity in any form

involving carcinogenicity, hepatotoxicity, nephrotoxicity, nor skin toxicity, as summarized in [Table 3](#).

#### ASO design

ASO design was performed using PFRED program which generated about 199 ASOs. Out of these, only 5 met the criteria of design mentioned in the methodology section. The ASOs are

presented in Table 4. Next, the top 5 ASO were checked for off-target with human transcriptome. Fortunately, all of them were above the set threshold (E-value < 0.5). This confirms the good design and selection of the program.

#### HFE prediction

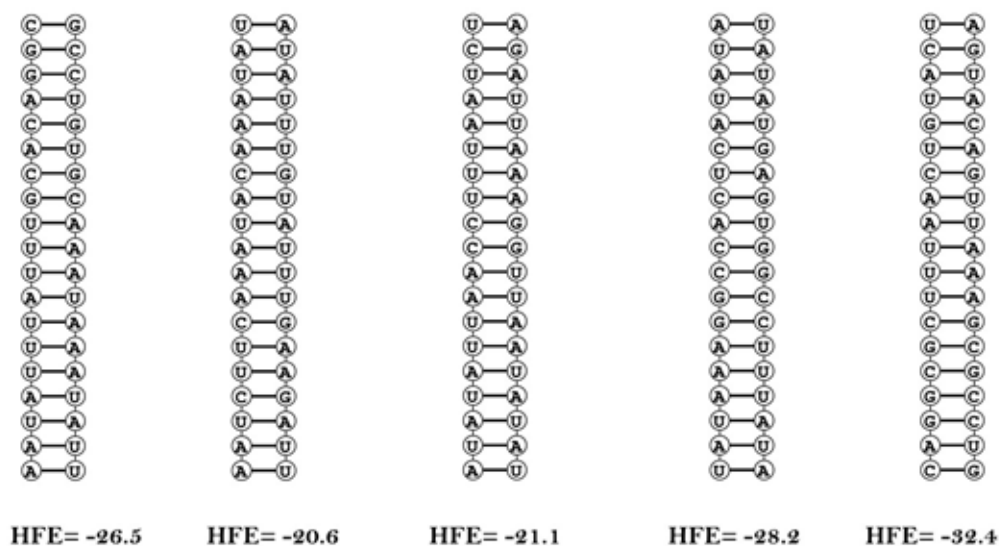
As demonstrated in Figure 4, ASO<sub>5</sub> followed by ASO<sub>4</sub> were the best in terms of complementarity strength to GR parent sense sequence giving HFE of -26.5 and -32.4 kcal/mol. However, ASO<sub>2</sub> and ASO<sub>3</sub> showed weak complementarity strength (-20.6 and -21.1) (Figure 4). This made ASO<sub>5</sub> as the best potential ASO to silence GR. To validate

those findings, RNA-RNA docking was made via HNADOCK platform.

The docking scores were inconsistent with DuplexFold output, ranking ASO1 as the best while ASO5 was the lowest (-365.68 vs. -218.64 kcal/mol). ASO<sub>2</sub> and ASO<sub>3</sub> were in-between (Figure 5). The difference in findings could be attributed to the fact that HFE is calculated based on sequence-based, bases hybridization whilst RNA-RNA docking is predicted according to structure-based complementation. In general, this leads us to the fact that all ASO are significantly potential GR silencers *in silico*, but the *in vitro* assays will give the most crucial result.

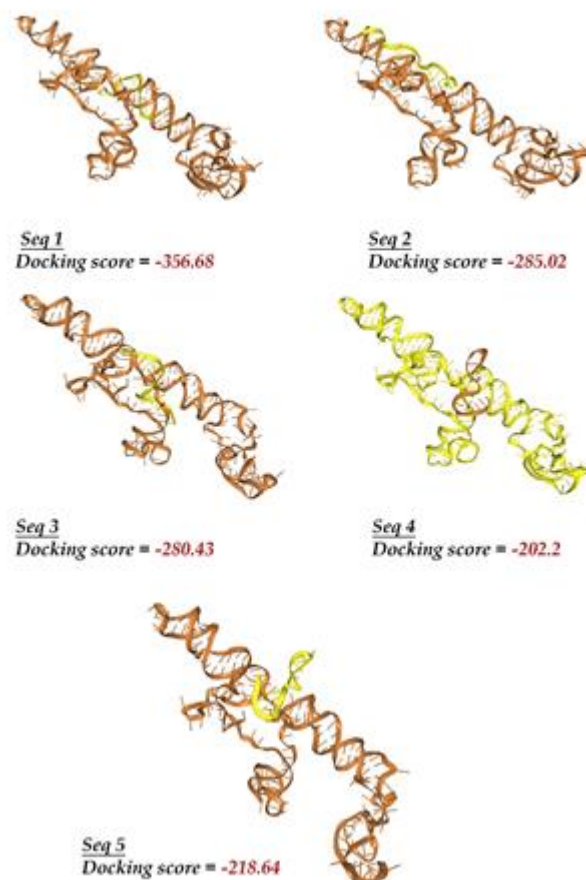
**Table 4.** Designed ASO along with Off-target E-value

No.	Parent sense oligo	Parent antisense oligo	Start	End	E-value
1	UAUAAAGGCCACUCAUAUA	UAUUGAGUGGCCUUUAUA	143	161	23
2	AUAUAUUAACCUUUAUCU	AGAUUAAAGGUUAAUAUAU	157	175	1.4
3	AAUCUUCAAAUACAAUAU	AUAUUUGUAUUUGAAGAUU	171	189	1.4
4	AAUAUUUAUUUGCACAGGC	GCCUGUGCAAUAAAUAUU	185	203	5.7
5	CAGGCGCUUUAACUGUACU	AGUACAGUUAAGCGCCUG	199	217	23



**Figure 4.** Sense-Antisense complementation and the predicted HFE.





**Figure 5.** RNA-RNA docking along with the docking score of each ASO-GR complex.

## Discussion

MRSA is a predator bacterial pathogen responsible for serious health issues and death cases each year particularly in USA and Europe. It still represents a medical obstacle since its discovery more than 60 years ago [28]. In fact, MRSA had emerged even before the introduction of methicillin into pharma market [29]. Riboswitches sense the concentration of a significant intracellular metabolite needed by an indispensable metabolic pathway in response to the environmental variation. Therefore, targeting bacterial riboswitches represent a new era for the development and testing novel drug candidate either of natural or synthesizable origins [30]. This instigates the medical community to search for novel target and novel

drugs to deal with such threat. Recent attention was directed toward MRSA riboswitch as a novel druggable target. Wang *et al.* [12] targeted FMN riboswitch with small molecules and proved their efficacy as potential drugs. Likewise, Traykovska and Penchovsky [31] designed and successfully applied ASO to silence S-adenosyl methionine-1 riboswitch of MRSA as a novel therapy. Theoretically, numerous studies reported the exploitation of bioinformatics approaches to provide unprecedented targets as well as inhibitors to the emerging pathogens including MRSA. This includes chalcone derivatives [32], benzamide derivatives [33], diverse inorganic coordination complexes [34], 2-Arylbenzimidazole Derivatives [35], and some natural flavonoids [36]. In this study, GR was the

target by small molecules (resveratrol derivatives) as well as ASO to inhibit the formed GR and prevent the transcription of GR. A dozen of resveratrol was docked into GR and proved their potency as potential natural inhibitors of such target. Besides, 5 ASO were designed against GR sequence and demonstrated their efficiency using HFE predicted by DuplexFold and docking to the native GR 3D structure by HNADOCK. GR serves as regulator of glucosamine synthase gene expression one of the contributing players to peptidoglycan biosynthesis. Upon binding its cognate ligand glucosamine-6-phosphate, a conformational change ensues leading to its activation and thus transformation to self-cleaving ribozyme. Ultimately, GR ribozyme activity cleaves glucosamine synthase mRNA in preparation to its decay by RNase J1 [37,38]. Recently, Mujwar and Pardasani [39] utilized computational methods to identify and virtually screen drug library against *Mycobacterium tuberculosis* riboswitches. Similarly, Anthrax riboswitches were also targeted as a means to inhibit and eradicate *Bacillus anthracis* bacterium *in silico* [40]. Furthermore, *Salmonella enterica* was another bacterium to be targeted via its riboswitch via a set of drug library *in silico* [41]. So, targeting such important genetic target to prevent MRSA membrane biosynthesis is an unprecedented aspect and the current study output deserves further validation *in vitro*.

## Conclusion

The present theoretical study elucidates the inhibition of Methicillin-resistant *Staphylococcus aureus* (MRSA) growth by a series of resveratrol derivatives, among which ZINC000100823225 emerges as the most promising potential inhibitor. This lead candidate exhibits a notable characteristic of adhering to of Lipinski's Rule of Five (Ro5) and demonstrates an absence of potential toxicity. Furthermore, I have designed and computationally demonstrated the efficacy

of five Antisense Oligonucleotides (ASO), which exhibit a significant capacity to silence the previously unexplored target, GR, *in silico*. This silencing potential is underscored by high hybridization energy and docking scores, as predicted by HNADOCK, while simultaneously ensuring the absence of any plausible off-target effects, particularly with respect to human genes. The combined utilization of these two strategic approaches holds substantial promise for mitigating the prevalence of MRSA and, as such, warrants further rigorous laboratory experimentation.

## List of abbreviations

ADMET	Absorption, distribution, metabolism, excretion, and toxicity
ASO	Antisense oligonucleotides
FMN	Flavin mononucleotide
GR	glms riboswitch
HFE	Hybrid free energy
MRSA	Methicillin-resistant <i>staphylococcus aureus</i>
VRSA	Vancomycin-resistant <i>S. aureus</i>
WHO	World Health Organization

## Ethics Approval and Consent to Participate

Not applicable.

## Human and Animal Rights

Not applicable.


## Conflict of interest

The author declares no known conflicts of interest.

## Disclosure statement

The authors declare that they have no conflict of interest.

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