Targeting RNA Polymerase Primary $\sigma^{70}$ as a Therapeutic Strategy against Methicillin-Resistant \textit{Staphylococcus aureus} by Antisense Peptide Nucleic Acid

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Abstract

Background: Methicillin-resistant \textit{Staphylococcus aureus} (MRSA) causes threatening infection-related mortality worldwide. Currently, spread of multi-drug resistance (MDR) MRSA limits therapeutic options and requires new approaches to “druggable” target discovery, as well as development of novel MRSA-active antibiotics. RNA polymerase primary $\sigma^{70}$ (encoded by gene \textit{rpoD}) is a highly conserved prokaryotic factor essential for transcription initiation in exponentially growing cells of diverse S. aureus, implying potential for antisense inhibition.

Methodology/Principal Findings: By synthesizing a serial of cell penetrating peptide conjugated peptide nucleic acids (PPNAs) based on software predicted parameters and further design optimization, we identified a target sequence (234 to 243 nt) within \textit{rpoD} mRNA conserved region 3.0 being more sensitive to antisense inhibition. A (KFF)$_3$K peptide conjugated 10-mer complementary PNA (PPNA2332) was developed for potent micromolar-range growth inhibitory effects against four pathogenic S. aureus strains with different resistance phenotypes, including clinical vancomycin-intermediate resistance S. aureus and MDR-MRSA isolates. PPNA2332 showed bacteriocidal antisense effect at 3.2 fold of MIC value against MRSA/VISA Mu50, and its sequence specificity was demonstrated in that PNA with scrambled PNA sequence (Scr PPNA2332) exhibited no growth inhibitory effect at higher concentrations. Also, PPNA2332 specifically interferes with \textit{rpoD} mRNA, inhibiting translation of its protein product $\sigma^{70}$ in a concentration-dependent manner. Full decay of mRNA and suppressed expression of $\sigma^{70}$ were observed for 40 $\mu$M or 12.5 $\mu$M PPNA2332 treatment, respectively, but not for 40 $\mu$M Scr PPNA2332 treatment in pure culture of MRSA/VISA Mu50 strain. PPNA2332 ($\geq$1 $\mu$M) essentially cleared lethal MRSA/VISA Mu50 infection in epithelial cell cultures, and eliminated viable bacterial cells in a time- and concentration-dependent manner, without showing any apparent toxicity at 10 $\mu$M.

Conclusions: The present result suggested that RNAP primary $\sigma^{70}$ is a very promising candidate target for developing novel antisense antibiotic to treat severe MRSA infections.

Introduction

As a frightening “superbug”, methicillin-resistant \textit{Staphylococcus aureus} (MRSA) has long been an overwhelming human pathogenic threat in healthcare-associated infections [1]. Its prevalence and adaptability in both community and hospital environment makes healthy patients and immune-deficient patients [2] at high risk of infection [3,4]. Its continued pathogenicity and virulence [5,6] causes invasive infection in bloodstream [7], essential organs, and tissues [8,9], therefore leads to severe clinical presentations and high mortality rate [4,10]. This is primarily due to the high incidence of methicillin-resistance that has failed almost all available antibiotics [11]. Furthermore, there has been an increase in reports of isolated MRSA strains developing multi-drug or vancomycin-(intermediate) resistance [12,13], which exacerbated antibiotic paucity. Interventions like vigilant monitoring of antibiotic susceptibilities and judicious use of culture-directed antibiotic agents have been a long-sought endeavor, yielding limited success [14]. Meanwhile, researchers and pharmaceutical industry have been driven to discover new MRSA-active agents (i.e. new chemical derivatives or compounds with new targets [15], virulence inhibitors [16], natural products, and vaccines [17]) and combination therapies, resulting in few ideal drugs or solutions [18]. Thus, antibacterial strategies that provide timely and effective therapeutic countermeasures are urgently required for possible outbreaks of MRSA infections. Particularly, specific RNA
silencing in bacteria by antisense antibacterial strategies can contribute to both aspects of the problem [19]. Antisense antibacterials are short (about 10- to 20-bases), synthetic DNA analogs that inhibit essential genes expression at mRNA level in a sequence-specific manner [20]. Therefore, antisense inhibition leads to bacteriocidal/bacteriostatic effect or restoration of bacterial susceptibility, which depends on the function of targeted gene. Synthetic antisense oligomers, especially peptide nucleic acid (PNA) [21] and phosphorodiamidate morpholino (PMO) [22], possess favorable properties in light of antisense antibacterial application, including improved targeting specificity, binding affinity, biological stability and access to a variety of chemical modification. Meanwhile, instead of simple mixture, cell penetrating peptides (CPP) can be covalently attached/conjugated at the end of PNA or PMO chain to enhance cellular uptake of antisense oligodeoxynucleotides (AS-ODNs) without affecting Waston-Crick base paring between antisense oligomers and targeted RNAs [23]. Synthetic peptide-PNA or peptide-PMO conjugates targeting growth-essential genes have shown to inhibit bacterial growth in pure culture and in infected tissue culture, Thus, a range of functional genes have been identified as potential targets [24]. However, only a few early reports provided preliminary proof-of-principle evidence on antisense targeting of S. aureus genes for growth inhibitory effect (i.e. peptide-PNA targeting lsbB [25], and phoB, lbwB, gycA, plus hmrB [26]) or restoration of antibiosis susceptibility (i.e. liposome-capsulated phosphorothioate oligodeoxynucleotides targeting mecA [27]) in pure culture. Targeting resistance mechanism in MRSA relies on elucidation of subtle intracellular self-regulation among related genes, the consequences of its antisense inhibition being too complicated to predict [28]. Thus, identification of growth-essential genes for more potent antisense inhibition in S. aureus would aid the development of new anti-MRSA agents [29].

Bacterial DNA-dependent RNA polymerase (RNAP) is a key enzyme in transcription regulation and gene expression. Its function requires coordination of a core enzyme (comprising five subunits α, β, β′ and ω) and an independent σ subunit that is reversibly recruited by core enzyme [30]. The RNAP core enzyme is responsible for transcription elongation, and different σs are in charge of transcription initiations from promoters that express genes in diverse function. The irreversible inhibition of RNAP thereby causes cell death. This has attracted much exploration for developing specific RNAP inhibitors (e.g. the rifamycins with fundamental clinical significance). The most developed σ70 family of σs, especially the primary σ70, is essential for initiating transcription of multiple genes in exponentially growing cells [31], which to our knowledge has not previously been demonstrated for antisense target validation in S. aureus. The primary σ’s are unique in structure, function and homology. The core regions of bacterial and eukaryotic RNAPs share structural and functional similarities, but the sequences of encoding genes are only partially homologous. Specifically, bacterial gene rpoD (encoding the primary σ70 of RNAP) shares the least homology in sequence with eukaryotic rpoD. Hence, in contrast to more conserved molecules, sequence-based drugs targeting rpoD products, including mRNAs, are less likely to cross react with host molecules. Most importantly, bacterial gene rpoD is highly conserved in identity and homologous in sequence among different pathogenic Staphylococcus species [32]. Such features are distinct advantages for developing narrow-spectrum anti-MRSA antisense agents [33].

In this study, by using four clinical pathogenic S. aureus genus with varying resistance patterns (including antibiotic sensitive, MRSA, MDR-MRSA and VISA), we report the identification of rpoD as a potent target for markedly bacteriocidal effect in vitro and ex vivo by antisense peptide-PNA conjugate.

Results

Transcript target site selection

The rpoD gene encoding bacterial RNAP primary σ70 is highly similar in sequence among S. aureus species. Sequence alignment of Staphylococcus rpoD by Blast showed 100% gene similarity in S. aureus, and other Staphylococcus (i.e. <85% in identity for S. epidermidis, S. lugdunensis, S. haemolyticus, S. pseudintermedius, and S. saprophyticus) (Table 1). Secondary structure of rpoD mRNA (Figure 1) was predicted by software RNA structure 4.6 and binding parameters (Table 2) were calculated by Oligo Walk plus PNALIGHT program. The combined data showed that each conserved region of rpoD mRNA has highly plausible sub-regions for antisense targeting. We selected total five plausible target sites for PNA synthesis (Table 2 & 3). All five mRNA regions are devoid of obvious stable secondary structures, thus they are theoretically accessible to complementary PNAs. All PNAs were covalently conjugated at the carbon terminus with peptide (KFF)6K (in which K is lysine and F is phenylalanine), acting as a carrier to facilitate delivery of PNA through stringent bacterial cell walls [34]. The efficacy of growth inhibition was determined by measuring minimal inhibitory concentration (MIC) in liquid bacterial culture. The results are illustrated for 4 S. aureus strains, including S. aureus ATCC29213, and clinical isolates of MRSA (i.e., Mu 50, WHO-2 and XJJING) (Table 3). In all strains, PNAs complementary to the rpoD mRNA nucleotides (nt) encoding 36 to 47, 159 to 170, and 312 to 323 (referring to anti-rpoD PNA 36, 159 and 312, respectively) were inactive at the highest concentration tested (25 μM); PNAs complementary to the rpoD mRNA nucleotides encoding 123 to 134 (anti-rpoD PPNA123) were equally active to S. aureus ATCC29213 and MRSA WHO-2, with a MIC value of 25 μM, whereas inactive to MRSA/VISA Mu50 and XJJING; PNAs complementary to the rpoD mRNA nucleotides encoding 233 to 244 (anti-rpoD PPNA233) were active to all 4 S. aureus strains, with a MIC value of 12.5, 12.5, 12.5 and 25 μM, respectively.

Anti-rpoD PPNA optimization

To optimize the design, we synthesized a set of antisense PPNAs based on anti-rpoD PPNA233 but differed in position of PNA and CPP, targeting site of PNA, PNA lengths, and CPP type (Table 3). Such design alterations should help to optimize the antimicrobial efficacy of obtained anti-rpoD PPNA233, and further illustrate the feasibility and sensitivity of selected target site in S. aureus rpoD mRNA. As shown in Table 3, the growth inhibitory effect of anti-rpoD PPNA2331 was abolished in all 4 S. aureus strains used above; anti-rpoD PPNA2332 and 2333 were potent, with increased activity against all 4 S. aureus strains. For anti-rpoD PPNA2332, the MICs were 12.5 μM, regardless of bacterial resistance phenotypes. Antisense specificity of anti-rpoD PPNA2332 was demonstrated in that PPNA with scrambled PNA sequence (Scr PPNA2332) showed no growth inhibition effect at the highest concentrations tested (40 and 25 μM, respectively). Anti-rpoD PPNA2333, which had the same PNA sequence of PPNA2332 but was attached to a considerably more effective peptide (RXX)6XB (in which R is arginine, X is 6-aminoacaproic acid and B is β-alanine), showed an equal MIC value of 6.25 μM among 4 S. aureus strains. However, the peptide-PNAs used in this study can be membrane active, and membrane permeabilization could result in nonspecific growth inhibition. This is the exact case for peptide (RXX)6XB, which showed undesirable growth inhibitory effect on...
all 4 S. aureus strains at 30 μM. Although this concentration was much higher than that of the conjugate PPNA2333 (MIC = 6.25 μM), it indicated that only anti-rpoD PPNA2332 provided improvements and properties that merit further evaluation.

Bacteriocidal antisense effect of anti-rpoD PPNA2332

The growth inhibitory effect of anti-rpoD PPNA2322 on bacterial cells were further examined by assessing cell growth and viability. In pure culture of MRSA Mu 50, we observed a time- and concentration-dependent inhibition of S. aureus growth in MH broth with PPNA2332. Scrambled PPNA2332 and control peptide (KFF)3K did not show any significant growth inhibitory effect at higher concentrations (Figure 2). There was no appreciable difference in growth between the cultures incubated with the Scr PPNA2332 or peptide (KFF)3K and those incubated without PPNA2332. All cultures grown without PPNA2332 increased by 4 log after 24 h of growth (1×10^6 CFUs/mL at 0 h) (Figure 3). However, PPNA2332 against rpoD mRNA was bactericidal in culture at ≥20 μM concentrations, with a >4 log reduction in CFUs at 24 h, compared with the culture at 0 h (P<0.01). Full elimination of live bacterial cells was observed for PPNA2332 at 40 μM when plated undiluted (limit of detection, 10 CFUs/mL). There were no difference in numerated colonies between all samples from cultures treated with Scr PPNA2332 or control peptide (KFF)3K and that of growth control. These results suggested that anti-rpoD PPNA2332 exerts a concentration-dependent bacteriocidal antisense effect.

Effect of anti-rpoD PPNA2332 on corresponding gene transcripts and protein expression

The reduction in viable bacterial cells we measured could be caused by a decay of specific transcripts of gene rpoD through antisense mechanism and subsequent decreased expression of protein σ70. To verify this possibility, we performed Reverse Transcription Polymerase Chain Reaction (RT-PCR) and western blotting for each individual transcript with total RNA and protein isolated from 18-hour MRSA/VISA Mu50 cultures upon inhibition of anti-rpoD PPNA 2332 at different concentrations. We observed that the levels of rpoD mRNA were greatly diminished in a concentration-dependent manner in treated cultures compared with untreated cultures (Figure 4A). Further, full suppression of rpoD mRNA was observed for PPNA 2332 treatment at 40 μM, which was probably due to steric blockage on top of mRNA decay effects. The level of the unrelated, constitutively expressed 16S rRNA gene transcript, assayed as a control RNA, did not change under any of the culture conditions. Again, the sequence specificity of the antisense gene knock-down effect was evaluated in each bacterial culture treated with Scr PPNA2332, and the expression of rpoD at mRNA level was not affected by the control condition at doses up to 40 μM (Figure 4A).

| Table 1. Homologies of gene rpoD among Staphylococcus species. |
|-------------------------------------------------------------|
| **Organism** | **GenBank** | **Similarity** | **Identity** |
|-----------------|-------------|----------------|--------------|
| *Staphylococcus aureus subsp. aureus* | | | |
| Mu50 | BA000017.4 | 100% | 100% |
| Mu3 | AP009324.1 | 100% | 100% |
| T0131 | CP002643.1 | 100% | 100% |
| Str. JKD6008 | CP002120.1 | 100% | 100% |
| TW20 | FN433596.1 | 100% | 100% |
| ED98 | CP001781.1 | 100% | 100% |
| JH1 | CP000736.1 | 100% | 100% |
| NCTC 8325 | CP000253.1 | 100% | 100% |
| USA300_TCH1516 | CP000730.1 | 100% | 100% |
| USA300_FPR3757 | CP000255.1 | 100% | 100% |
| ST398 | AM990992.1 | 100% | 100% |
| MRSA252 | BX571856.1 | 100% | 100% |
| MSSA476 | BX571857.1 | 100% | 99% |
| JKD6159 | CP002114.2 | 100% | 99% |
| ED133 | CP001996.1 | 100% | 99% |
| **Other Staphylococcus** | | | |
| Staphylococcus epidermidis RP62A | CP000297.1 | 100% | 85% |
| ATCC 12228 | AE015929.1 | 100% | 85% |
| Staphylococcus lugdunensis HKU09-01 | CP001841.1 | 100% | 84% |
| N920143 | FR870271.1 | 100% | 84% |
| Staphylococcus haemolyticus JCSC1435 | AP008716.1 | 100% | 84% |
| Staphylococcus carnosus subsp. carnosus TM300 | AM295250.1 | 100% | 84% |
| Staphylococcus pseudintermedius ED99 | CP002478.1 | 100% | 83% |
| HKU10-03 | CP002439.1 | 100% | 83% |
| Staphylococcus saprophyticus subsp. saprophyticus ATCC 15305 | AP008934.1 | 98% | 83% |

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To ascertain whether the antisense inhibition of rpoD at mRNA level could lead to decreased expression of its protein product, the expression of σ70 was tested. Interestingly, inhibiting the rpoD mRNA by PPNA2332 led to decreased expression of σ70 in a concentration-dependent manner (Figure 4B). But the Scr PPNA2332 treatment showed no effect on the expression of σ70. These results demonstrated that RNAP primary σ70 was crucial for sustaining bacteria surviving, and knock-down of rpoD by PPNA2332 induced transcript decay and unsuccessful translation of transcripts.

Protective effect of anti-rpoD PPNA2332 on human cultured cells from single MRSA infection

To examine the antibacterial potential of anti-rpoD PPNA2332 in the presence of eukaryotic cells as well as to evaluate its possible toxicity, we tested it against single clinical MRSA/VISA strain Mu50 grown in gastric mucosa originated epithelial cell (GEP) culture medium. In this medium, which is likely to be more representative of in vivo application, PPNA2332 was >10-fold more potent (MIC = 1 μM) than in MH broth. Furthermore, epithelial cell cultures were artificially infected with 10⁷ CFU/mL of invasive MRSA/VISA Mu50 cells, and anti-rpoD PPNA2332, Scr PPNA2332, and control peptide (KIF)₃K were added immediately post-infection or tissue culture medium only. This system can be viewed as a very simple model for the growth of an extracellular pathogen in a host. Bacterial CFUs from each condition were measured at 2, 6, and 24 h after infection. MRSA/VISA Mu50 grew 2 log in 24 h in the presence of medium alone (Figure 5). At 2 h after infection, PPNA2332 in infected GEP had reduced bacterial CFUs by 0.65 log (Figure 5). Further reduction of CFUs was continued over time, and by 6 h PPNA2332 caused fully elimination of CFUs, compared with infected GEP without PPNA treatment. The difference in killing was most pronounced after 24 h, when there was a 7.45 log difference (P<0.01) in CFUs between bacterial cells in GEP and the addition of PPNA2332 (Figure 5). The addition of the controls (Scr PPNA or peptide) had no appreciable effect on antisense killing of bacterial cells, compared with bacterial cells grown in GEP alone.

Figure 1. Secondary structure of E. coli RNA polymerase primary σ70 mRNA. The target region of the 10-mer peptide nucleic acid (nucleotides 234–243) is indicated by the bold line.

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These results were confirmed by microscopic examination of the cultures, which showed that anti-rpoD PPNA2332 treated GEP cultures appeared the same as uninfected cultures (Figure 6). All infected controls appeared the same, independent of treatment, and showed few GEP cells but massive amounts of bacteria, which made the cultures appear turbid. Results obtained using Scr PPNA2332 and peptide (KFF)3K were similar to those obtained in MRSA/VISA Mu50 cells in GEP culture only (data not shown).

### Table 2. Binding parameters predicted by software Oligo walk 5.0 and PNALIGHT.

| PNA sequence                  | Conserved region of rpoD | Target Site* | Parametersb | DNA:PNA | GC (%) | overall ΔG | Duplex ΔG | Oligo-self ΔG | Oligo-oligo ΔG | Tm[°C]d |
|-------------------------------|--------------------------|--------------|-------------|---------|--------|------------|-----------|---------------|---------------|---------|
| 5'-gtcgcagaattc-3'             | r1.1                     | 36–47        | 41.7        | −16.2   | −16.6  | 0          | −5.2      | 60.0          |
| 5'-agtgggatcga-3'              | r1.2                     | 123–133      | 33.3        | −14.5   | −14.9  | 0          | −2.7      | 56.9          |
| 5'-ctacatcaggta-3'             | r1.2                     | 159–170      | 33.3        | −15.3   | −15.7  | 0          | −2.5      | 53.4          |
| 5'-ttttctgctagc-3'             | r3                      | 233–244      | 58.3        | −17.6   | −18    | 0          | 0         | 65.1          |
| 5'-tttctgctagc-3'              | r3                      | 233–242      | 40          | −a      | −       | −          | −         | 43.0          |
| 5'-ttttctgctacg-3'             | r3                      | 234–243      | 40          | −       | −       | −          | −         | 45.9          |
| 5'-cgcccaattct-3'              | r4                      | 312–323      | 50          | −17.8   | −18.2  | 0          | −1.5      | 57.5          |

*a*Numbering from the first base of the gene rpoD;

ΔG means free energy; index for each parameter: GC%=60%, overall ΔG<−10 kcal/mol, Duplex ΔG<−25 kcal/mol, oligo-self ΔG<−1.1 kcal/mol, oligo-oligo ΔG<−8 kcal/mol, Tm≥50 °C.

bCalculated melting temperature in °C of matching PNA/DNA hybrids with no dangling ends.

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### Table 3. MIC of anti-rpoD peptide-PNAs for quality control and clinical strains of Staphylococcus aureus in M-H broth culture.

| PNA designation and sequencea | No. of PNA bases | MICb for S. aureus strains (µM) | ATCC MRSA/VISA/MDR | WHO-2 | XIJING |
|-------------------------------|------------------|---------------------------------|---------------------|-------|--------|
| anti-rpoD PPNA (without spacer between PNA and peptide) |  |  |  |  |  |
| 36 5'-gtcgcagaattc-(KFF)3K-3' | 12 | >25 | >25 | >25 | >25 |
| 123 5'-agtgggatcga-(KFF)3K-3' | 12 | 25 | >25 | 25 | 25 |
| 159 5'-ctacatcaggta-(KFF)3K-3' | 12 | >25 | >25 | >25 | >25 |
| 233 5'-ttttctgctagc-(KFF)3K-3' | 12 | 12.5 | 12.5 | 12.5 | 25 |
| 312 5'-cgcccaattct-(KFF)3K-3' | 12 | >25 | >25 | >25 | >25 |
| anti-rpoD PPNA based on 233 (with spacer between PNA and peptide) |  |  |  |  |  |
| 2331 5'-t(KFF)3KXKX (KFF)3KX-3' | 10 | >25 | >25 | >25 | >25 |
| 2332 5'-t(KFF)3KXKX (KFF)3KX-3' | 10 | 12.5 | 12.5 | 12.5 | 12.5 |
| 2333 5'-t(RKK)3KXKX (KFF)3KX-3' | 10 | 6.25 | 6.25 | 6.25 | 6.25 |
| Scr 2332 5'-t(KFF)3KXKX (KFF)3KX-3' | 10 | >40 | >40 | >40 | >40 |
| Scr 2332 5'-t(RKK)3KXKX (KFF)3KX-3' | 10 | >25 | >25 | >25 | >25 |
| Free peptides (KFF)3KXKX | >60 | >60 | >60 | >60 | >60 |
| RXRXXXRXRXXKX | 30 | 30 | 30 | 30 | |
| Controls |  |  |  |  |  |
| oxacillin | 0.5 | >1024 | 512 | >1024 |  |
| cefazidine | 8 | 1024 | 256 | 256 |  |
| ampicillin | 1 | >1024 | 256 | >1024 |  |

a*rpoD*, RNA polymerase sigma 70; PNA, peptide nucleic acid; PPNA means peptide conjugated PNA; The PNAs are written from their N to their C terminus, and the N terminus corresponds to the 5' end of a conventional oligonucleotide; “K” indicates lysine, F indicates phenylalanine, “X” indicates 6-aminohexanoic acid, “B” indicates p-alanine, and eg1 indicates glycine; Scr means PNA with a scrambled base sequence (as control);

bMinimal inhibitory concentrations (MIC) were the lowest PNA concentrations that prevented bacterial growth by visual inspection after overnight (24 h) growth from an inoculum of 105 CFU/mL. “VISA” is abbreviation for vancomycin-intermediate resistance Staphylococcus aureus; “MRSA” is abbreviation for methicillin-resistant Staphylococcus aureus; XIJING means clinical MRSA isolate from patients in Fourth Military Medical University affiliated XIJING hospital.

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suggest that the 234–243 nt of pathogenic MRSA. By using peptide-PNA conjugates, our results implied that targeting rpoD mRNA (in conserved region 3) is most sensitive to antisense inhibition. An optimized (RXR)4XB conjugated PNA complementary to this sequence exerted by far the most potent (MIC = 6.25 μM) antisense growth inhibitory effect against clinical isolates of pathogenic MRSA, which implied that targeting rpoD by high concentration of PPNA would lead to complete decay of rpoD mRNA and finally concomitant cell death. More importantly, micromolar PPNA2332 treatment was able to significantly rescue 100% of MRSA-infected epithelial cells, with full elimination of MRSA cells in co-culture medium. This highly implied the therapeutic potential of PPNA2332 in the treatment of MRSA infections by targeting rpoD.

A challenging aspect of identifying essential genes in bacteria for antisense inhibition mainly involves efforts to locate the exact

**Discussion**

Rapid evolution of resistance genes developing among Gram-positive pathogens has unfortunately exceeded our ability to develop new antimicrobials [35]. The methicillin-resistance, vancomycin-intermediate resistance or MDR- S. aureus present formidable challenges for traditional antibiotic discovery [36]. Novel antisense antibacterial strategies have been introduced and can efficiently provide potential drugs to combat emerging or re-emerging pathogens.

The conserved bacterial DNA-dependent RNA Polymerase (RNAP), is the target of rifamycins and is of great interest as a potential target for developing antisense bacteriocidal agents. Thus, we set out to target rpoD (encoding RNAP primary σ70) in S. aureus by antisense strategy, assuming it was essential for survival of pathogenic MRSA. By using peptide-PNA conjugates, our results suggest that the 234–243 nt of rpoD mRNA (in conserved region 3) is most sensitive to antisense inhibition. An optimized (RXR)4XB conjugated PNA complementary to this sequence exerted by far the most potent (MIC = 6.25 μM) antisense growth inhibitory effect against clinical isolates of pathogenic MRSA in vitro. Our study also showed that antisense targeting of rpoD by PPNA2332 could significantly inhibit growth of S. aureus genus of different resistance phenotypes. The effect of PPNA2332 was sequence specific, because a scrambled-sequence PNA did not inhibit growth of S. aureus genus, nor did incubation with the peptide (KFF)3K alone. In vitro gene-specific bacteriocidal activity was also seen in clinical isolate of MRSA, which implied that targeting rpoD by high concentration of PPNA would lead to complete decay of rpoD mRNA and finally concomitant cell death. More importantly, micromolar PPNA2332 treatment was able to significantly rescue 100% of MRSA-infected epithelial cells, with full elimination of MRSA cells in co-culture medium. This highly implied the

**Figure 2. Effects of anti-rpoD PPNA2332 on the growth of MRSA/VISA Mu50 in pure culture.** Anti-rpoD PPNA2332 was added to cell cultures containing 1.0×10^5 CFU/mL MRSA/VISA Mu50 to a final concentration of 6.25, 12.5, or 25 M. Additional cell cultures were treated with free MH broth, scrambled PPNA2332 (final concentration of 40 μM), and peptide (KFF)3K (final concentration of 40 μM) in a volume equal to that of the PPNA2332 preparation as controls. The growth of different groups of MRSA/VISA Mu50 cells was monitored by using OD measurements. The data are shown as means for 2 samples from 2 independent tests.

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**Figure 3. Bacteriocidal effects of anti-rpoD PPNA2332 on the viable cells of MRSA/VISA Mu50 in pure culture.** Anti-rpoD PPNA2332 was added to cell cultures containing 1.0×10^5 CFU/mL MRSA/VISA Mu50 to a final concentration of 5, 10, 20, or 40 μM. Additional cell cultures were treated with free MH broth, scrambled PPNA2332 (final concentration of 40 μM), and peptide (KFF)3K (final concentration of 40 μM) in a volume equal to that of the PPNA2332 preparation as controls. Aliquots of each culture were collected at 0 h and 24 h, diluted, and inoculated onto solid MH agar. The number of CFU was calculated from the number of colonies growing on plates. The 24 h counts for 40 μM PPNA2332 were 0 CFUs/mL. The data are shown as means ± SD for 2 samples from 2 independent test. *, P<0.01 for comparison to control values.

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**Figure 4. Specific inhibition of RNAP primary σ70 gene transcript and protein expression by anti-rpoD PPNA 2332.** Triplicate bacterial cultures were grown for 18 h in 100 μL of MH broth in the presence of anti-rpoD PPNA2332 and scrambled PPNA2332 at different concentrations with MRSA/VISA Mu50 cells, respectively. The anti-rpoD PPNA2332 was added once at the start of the growth period. After 18 h of growth, total RNA and protein was isolated from treated and untreated cultures. (A) Cellular levels of σ70 RNA were determined by RT-PCR. The reduction in the amount of RT-PCR product corresponding to gene rpoD in cells treated with anti-rpoD PPNA2332 at different concentrations was determined as the product specific for rpoD RNA relative to products for 16s rRNA. (B) Expressions of σ70 protein in different treatment were analyzed by western blotting and were quantitated densitometrically.

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targeting site within gene bases for realizing potent and specific antisense inhibitory effect of complementary AS-ODNs. Proof-of-principle evidence from previous study on identifying genes essential for growth in \textit{S. aureus} was first offered by Nekhotiaeva et al. \cite{26}. They have used four endogenous genes in \textit{S. aureus} RN4220, including \textit{phoB} (encoding alkaline phosphatase), \textit{fmhB} (involved in cell wall biosynthesis), \textit{gyrA} (involved in DNA replication), and \textit{hmrB} (an ortholog of the \textit{E. coli} \textit{acpP} gene and highly sensitive to antisense inhibition in \textit{E. coli}). And they tested the bacterial growth inhibition efficacy of antisense peptide-PNAs to verify the essentiality of these genes. They only designed 2 or 3 peptide-PNAs targeting the start codon region and upstream Shine-Dalgarno (SD) region in each gene, which were previously reported for higher sensitivity to antisense inhibition. In all cases, they observed concentration-dependent growth inhibition with sequence-specificity. However, MIC values were not determined and the full-course inhibition of bacterial growth was only observed for peptide-PNA targeting \textit{fmhB} at the concentration of 10 nM. Limited information from two other studies have shown similar MIC values (15 nM) for AS-ODNs targeting \textit{fabI} \cite{25} or \textit{adk} \cite{37} in \textit{S. aureus} RN4220, regardless of ODN type (i.e., DNA or PNA) and delivery strategies (i.e., none or peptide mediated).

Bacterial primary $\sigma^{70}$'s are responsible for transcriptional initiation of multiple essential genes that play important roles on cell survival and proliferation in exponential phase. Bacteria have no bypass mechanism to retrieve the sacrifice for the inactivation of primary $\sigma^{70}$'s, which certainly leads to transcription failure and subsequent loss of functional proteins. Common pathogenic \textit{S. aureus} share highly homologous sequences in gene \textit{rpoD} (Table 1). However, little evidence has been provided to identify the accessible and sensitive regions for potent antisense inhibition. The inhibition profile from our preliminary target site selection results clearly indicates that the conserved region 3 of \textit{rpoD} containing 233 to 244 nt sequence is of most sensitivity among five selected regions (Table 2&3). More potent growth-inhibition effect was obtained only for optimized PNA encompassing the 234 to 243 nt sequence. This further confirmed the least indispensable target sequence for effective antisense inhibition. Notably, the growth inhibition effect of the 10-mer \textit{rpoD}-targeting PNA is through antisense mechanism, which was reflected by the concentration-dependent decay of \textit{rpoD} mRNA transcript and decreased protein expression of $\sigma^{70}$ observed for \textit{PPNA2332} treated bacterial cells rather than those treated by Scr \textit{PPNA2332}.

Certain peptide-PNA and peptide-PMO oligomers targeting growth-essential genes are more bacteriocidal \textit{in vitro} relative to molar or mass equivalent doses of ampicillin and rifampicin, and they act with gene and sequence selectivity. Potent bacteriocidal antisense effect of anti-\textit{rpoD} \textit{PPNA2332} was highly presumable to resemble pharmacological aspects (or mode of action) of rifampicins, the famous broad-spectrum RNAP inhibitors that accumulate in bacterial cells to exert post antibiotic effect, with comparatively higher MICs than other types of antibiotics \cite{38}.

*Figure 5. Protective effect of anti-\textit{rpoD} \textit{PPNA2332} on epithelial cell culture infected with noninvasive single MRSA/VISA Mu50 infection.* The symbols represent time points of cell harvest (2, 6, and 24 h) and are displayed as the mean number of colony-forming units (CFUs) per milliliter. Error bar represents SD. The 6 h counts for 1 nM \textit{PPNA2332} were 0 CFUs/mL. For all observations in co-cultures (GEP+MRSA/VISA Mu50) treated, with the limit of detection indicated as 10 CFUs/mL. Scr \textit{PPNA2332}, Scrambled sequence control \textit{PPNA2332}; GEP, gastric mucosa originated epithelial cell.
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*Figure 6. Light micrographs of epithelial cell cultures.* The images show epithelial cell cultures grown in DMEM/HIGH Glucose, 10% FCS. The column at left shows epithelial cell cultures without added MRSA/VISA Mu50; the right column shows epithelial cell cultures that were inoculated with MRSA/VISA Mu50. The top panels of each column show cultures not treated with PPNA, and the rows below show cultures treated with increasing amounts of anti-\textit{rpoD} \textit{PPNA2332} (1, 5, and 10 nM). Magnification, $\times$100.
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we firstly demonstrate that bacterial animal and human toxicity, and pharmacokinetic behavior of
of PPNAs were observed to eliminate bacteria dramatically in a broth. However, accumulation and sustained bacteriocidal effect of PPNA2332 was demonstrated by the increase of CFUs in four S. aureus strains with different characteristics in resistance phenotype. All PNAS used are covalently conjugated with the same cell penetrating peptide (KFF)3K at the carbon terminus (corresponding to the 3’ end of a conventional oligonucleotide).

To further optimize the anti-\(\sigma^D\) PNA that showed best antisense inhibitory efficacy in preliminary target site selection, de novo design in ways of (i) adding glycine spacer between PNA and CPP; (ii) shortening PNA length; and (iii) utilization of newly developed CPP (RXR)\(_2\)XB for more efficient PNA delivery into larger scale of bacterial species, were performed. Optimal anti-\(\sigma^D\) PNA was selected by minimal inhibitory concentration (MIC) results determined for clinical isolates of pathogenic S. aureus species.

Materials and Methods

Chemicals
All antibiotics used were purchased from the National Institute for the Control of Pharmaceutical and Biological Products (Beijing, China). All other chemicals and solvents were of analytical grade.

Bacterial strains, CPP, and PNA
The S. aureus strain ATCC29213 (antibiotic sensitive, for quality control) and WHO-2 (MRSA) were obtained from the Chinese National Center for Surveillance of Antimicrobial Resistance (Beijing, China), and clinical strain MRSA Xijing (MDR) was isolated from cultures of sputum or catheter samples from patients in Xijing Hospital (Xi’an, China). MRSA/VISA Mu50 (ATCC700699) was purchased from MicroBiotics (Minnesota, USA). All strains expressed \(\sigma^D\), which was confirmed by PCR detection (data not shown). CPPs were synthesized and purified at Genotide, Inc. (Xi’an, Shanxi, China). PNNAs were synthesized and purified at Panagene Inc (DAEHEON, Korea). peptide-conjugated PNNAs were synthesized by manual coupling chemistry and purified at Panagene, Inc (DAEHEON, Korea), with the base and amino acid sequences shown in Table 3.

PNA target site selection and anti-\(\sigma^D\) PNA (PPNA) optimization
Sequence alignment of gene \(\sigma^D\) was performed to decide the region that showed the highest sequential homology among different S. aureus species. Secondary structure of \(\sigma^D\) mRNA predicted by RNA structure 4.6 software. DNA:RNA and DNA:DNA paring parameters referenced by Oligo Walk 3.0 and PNALIGHT program were used to decide the site that showed best binding affinity. Building from evidence that 12-base antisense PNAS was an effective length [23], five plausible target sites located in the region that satisfied both above aspects were chosen for synthesis of corresponding antisense PNAS at 12-base length. Effective target site in \(\sigma^D\) was verified by modified MIC test of PPNA in four S. aureus strains with different characteristics in resistance phenotype. All PNAS used are covalently conjugated with the same cell penetrating peptide (KFF)3K at the carbon terminus (corresponding to the 3’ end of a conventional oligonucleotide).

To further optimize the anti-\(\sigma^D\) PNA that showed best antisense inhibitory efficacy in preliminary target site selection, de novo design in ways of (i) adding glycine spacer between PNA and CPP; (ii) shortening PNA length; and (iii) utilization of newly developed CPP (RXR)\(_2\)XB for more efficient PNA delivery into larger scale of bacterial species, were performed. Optimal anti-\(\sigma^D\) PNA was selected by minimal inhibitory concentration (MIC) results determined for clinical isolates of pathogenic S. aureus species.

Bacterial growth assay and susceptibility testing
MICs were determined twice by the microdilution assay in sterilized 96-well polystyrene microtiter plates according to the broth microdilution guideline based on Good et al. [23]. Briefly, single colony overnight culture of subjected bacterial cells was diluted to give a final inoculation at \(10^5\) CFU/mL in duplicate 100 \(\mu\)L Mueller Hinton (MH) broth in a low-binding 96-well microtiter plate. Stocks of PPNA, Scr PPNA and CPPs were added immediately to the indicated final concentrations. Plates were incubated at 35°C in aerobic environment for 24 h of overnight culture. MICs were the lowest PNA concentrations that prevented bacterial growth by visual inspection after overnight (24 h) growth. To verify the specific antisense effect, peptide was also conjugated to a scrambled PNA with a base sequence that was not complimentary to the selected target site of \(\sigma^D\) mRNA. To control for nonspecific toxicity, the MIC of the control CPPs (KFF)3K and (RXR)\(_2\)XB in all cases, was above the limit of measurement, which was 60 or 30 \(\mu\)M.

To determine the growth curve for MRSA/VISA Mu50 in the broth medium, cells were diluted and mixed with no PNA, anti-\(\sigma^D\) PPNAs 2332 (6.25, 12.5, and 25 \(\mu\)M), and scrambled PPNAs 2332 (40 \(\mu\)M). The growth rate of the cells was monitored by measuring the OD values at 620 nm with a microplate reader (Bio-Rad Laboratories, Tokyo, Japan) at different time points (0, 2, 4, 6, 8, 10, 12, 14, 16, 18, and 24 h). After 24 h culture, viable cells were determined by plating 50 \(\mu\)L samples from each wells onto Mueller-Hinton agar plates in appropriate diluted concentrations, and numbers of colonies was counted.
Reverse transcript (RT)-PCR

To compare \( \sigma^{32} \) RNA levels in inhibited and uninhibited bacterial cells, duplicate subjected MRSA/VISA Mu50 were grown as described above and treated with either no PPNA inhibitor, different concentrations of anti-\( \sigma^{32} \) PPNA2332, or scrambled PPNA2332 for 18 h. Cell amounts were verified by plating. Total RNA from each culture was prepared using the RNeasy Mini Kit (QIAGEN China Co. Ltd, Shanghai, China) or RNAprep pure Cell/Bacteria Kit (TIANGEN BIOTECH Co, Ltd, Beijing, China), and followed by reverse transcript using PrimeScript RT reagent Kit with DNA Eraser (TAKARA BIO INC, Kyoto, Japan). PCR was performed with the Premix Taq RT-PCR System (TAKARA BIO INC, Kyoto, Japan) according to the manufacturer’s instructions. Primers specific for \( \sigma^{32} \) gene in subjected \( S. aureus \) were listed as followed. Primers specific for MRSA/VISA Mu50 \( \sigma^{32} \) were 5’-CAGATACGTGACGAGAAA-3’ and 5’-GAAATAATACCAAGAACAG-3’; Primers specific for MRSA/VISA Mu50 16S rRNA were 5’-CGTTGGAATACCTATAAAGACT-3 and 5’-GATTCCCTACTGCTGCTG-3’. The PCR fragment encompasses the targeting site of PPNA2332 in \( \sigma^{32} \) gene. Amplification was performed in a Gradient thermal cycler (BioRad laboratories Inc., Hercules, CA, USA) under the following condition: denaturation at 95°C for 3 min for the first cycle and for 30 s thereafter, annealing at 55°C for 30 s, and extension at 72°C for 40 s for 32 repetitive cycles. Final extension was at 72°C for 10 min. 16S rRNA was used as an internal control. The PCR products were analyzed by electrophoresis on a 1% agarose gel.

Western blotting

To compare \( \sigma^{32} \) expression levels in inhibited and uninhibited bacterial cells, duplicate subjected MRSA/VISA Mu50 were grown as described above and treated with either no PPNA inhibitor, different concentrations of anti-\( \sigma^{32} \) PPNA2332 and scrambled PPNA2332 for 18 h. Anti-\( \sigma^{32} \) PPNA treated bacterial cells and untreated controls were lysed using lysis buffer (Dingguo Biotech Co, Ltd, Beijing, China) containing lysozyme (100 g/mL) and PMSF (1 mM). Cell lysates were quantitated, resolved, blotted and visualized essentially as previously described. Equal amounts of protein were loaded and separated on 12% SDS-polyacrylamide gel and were then transferred to polyvinylidene difluoride membranes (Millipore Corporation, Billerica, MA, USA). The bacterial RNAP \( \sigma^{32} \) monoclonal antibody (Abcam, Cambridge, MA, USA) was used at dilution of 1:1000 for overnight blotting at 4°C; the secondary horseradish peroxidase-conjugated anti-mouse antibody was used at dilution of 1:2000.

Epithelial cell culture, single bacterial infection, and anti-\( \sigma^{32} \) PPNA2332 treatment

Epithelial cells (gastric mucosa originated, a gift from Dr. Na Chai) were plated in 96-well culture dishes (Falcon, Franklin Lakes, NJ) at a concentration of 1.5 x 10^5 cells/mL in a volume of 0.200 µL and grown at 37°C for 48 h in Dulbecco’s minimal essential medium (high sugar DMEM, HyClone, Logan, Utah) supplemented with 10% fetal bovine serum (HyClone, Logan, Utah) in 95% air-5% CO₂. An overnight respective culture of MRSA/VISA Mu50 (ATCC 700699) was diluted by high sugar DMEM with 10% fetal bovine serum for fixed-concentration experiments, and transferred to wells containing epithelial cells. The starting bacterial inoculum for single infection was 1 x 10^7 CFU/mL in a 100 µL volume. Anti-\( \sigma^{32} \) PPNA 2332 (1, 5, and 10 µM) and scrambled PPNA2332 (20 µM) were immediately added, and the cultures were incubated at 37°C in 95% air-5% CO₂ for 24 h. After 24 h, cultures were examined and photographed with a Nikon inverted light microscope under x100 magnification. And culture supernatant was removed, diluted to appropriated concentration and plated on MH agar to measure viable bacterial cells.

Statistical Analysis

Values shown in the graph are means of two or three replicates from independent experiments. Results are expressed as mean or mean ± SD where indicated. For the in vitro experiments, cell viability (in units of CFUs per milliliter) at 24 h was calculated for MRSA/VISA Mu50 and PPNA2332 treatment condition, including no treatment. The difference (in units of log count +0.5) between cultures treated with PPNA2332 and untreated cultures was analyzed by means of the paired Student t test. Values were linearly transformed by the addition of a constant, 0.5, to allow statistical testing of the log counts, as samples were completely sterilized by PPNA2332 (i.e., 0 CFUs/mL). A probability value of P<0.01 was considered indicative of statistical significance.

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Author Contributions

Conceived and designed the experiments: HB XXL. Performed the experiments: HB GJS YY ZH XYX YZ JRM. Analyzed the data: HB GJS YY. Contributed reagents/materials/analysis tools: HB GJS YY ZH XYX YZ JRM. Wrote the paper: HB XXL.

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