Antimicrobial Resistance: Techniques to Fight AMR in Bacteria – A Review

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Abstract

Antimicrobials or antibiotics were the important revelations of the last century, however, it came along with a silent curse that people care less to talk about. Antimicrobial resistance (AMR) which emerged alongside antibiotics in the last century has been a significant concern for scientists and policymakers. Since their discovery, it has been noted that the widespread use of antibiotics is the primary cause of bacteria developing antimicrobial drug resistance. Despite the recognition of this issue, it is challenging to curtail the widespread use of antibiotics because they are essential for treating various infections. Paradoxically, the necessity of using these drugs becomes an inadvertent advantage for bacteria to evolve resistance mechanisms. This dilemma creates a seeming stalemate in our battle against these tiny microorganisms. Delaying action could have dire consequences, potentially leading to the emergence of stronger superbugs that pose a serious threat to the entire human population. The recent COVID-19 pandemic serves as a stark reminder of the devastating impact a small microbe can have on global health. This paper delves into the mechanisms of antimicrobial resistance in bacteria, the evolution of superbugs and the innovative techniques employed by scientists to combat these challenges. Taking proactive steps is crucial to avoid a future where we are at the mercy of increasingly resilient microbes.

Keywords: Antimicrobial Resistance, Superbugs, Efflux pumps, β-Lactamases, Nanoparticles, CRISPR/Cas9

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INTRODUCTION

Antibiotics were systematically thought to be the important discovery of the previous century. Their boundless use transformed healthcare after their debut and the names of scientists like Paul Ehrlich and Sir Alexander Fleming who introduced them to us are still discussed. Various antibiotics have effectively controlled infectious diseases that were once untreatable and fatal, thus preventing millions of deaths each year. However, due to the overuse and misuse of antibiotics, the effectiveness of these drugs is now threatened by the rise of antibiotic resistant bacteria. Fleming, in his days, had noted the insensitive use of these drugs might result in the development of bacterial strains which might become immune. This ability of microorganisms, such as bacteria, viruses and some parasites, to develop resistance to antimicrobial drugs has been termed antimicrobial resistance (AMR).

The World Health Organization (WHO) has recognized AMR as one of the ten major worldwide public health challenges. It is a significant global concern that requires coordinated efforts to combat its spread and ensure the continued effectiveness of these vital drugs. This has prompted researchers and pharmaceutical companies to intensify their efforts in drug research and development, with a focus on understanding resistance mechanisms and developing novel antibiotics and alternative treatment strategies. In this article, we have explored the various AMR mechanisms and discussed methods to prevent it.

Antimicrobials

Antimicrobials are a diverse group of substances that are pivotal in the fight against infectious diseases. These compounds, including antibiotics, antivirals, antifungals and antiparasitic, work by either killing or inhibiting the growth of microorganisms such as bacteria, viruses, fungi and parasites. They play a crucial role in medicine, agriculture and veterinary care, combating infections in humans, animals and plants. Antimicrobials encompass a wide array of compounds produced through various means, including natural sources like bacteria, fungi and plants, as well as synthetic processes in
Table 1. Overview of Antibiotic Resistance Mechanism Employed by Bacteria

<table>
<thead>
<tr>
<th>Antibiotic</th>
<th>Target</th>
<th>Mechanism Of Action</th>
<th>Mechanism Of Resistance</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>β-Lactams</td>
<td>β-Lactamases that hydrolyse the β-lactam ring.</td>
<td>Production of β-lactamases that hydrolyse the β-lactam ring.</td>
<td>Alteration or decrease in expression of PBPs.</td>
<td>59</td>
</tr>
<tr>
<td>Vancomycin</td>
<td>D-Ala-D-Ala terminus of peptidoglycan precursors</td>
<td>Converting of D-Ala-D-Ala to D-Ala-D-Lac or D-Ala-D-Ser</td>
<td>Modification of target site through methylation.</td>
<td>60</td>
</tr>
<tr>
<td>Macrolides</td>
<td>Bacterial ribosomes</td>
<td>Modification of target site through methylation.</td>
<td>Active efflux of antibiotic.</td>
<td>61</td>
</tr>
<tr>
<td>Tetracyclines</td>
<td>Bacterial ribosomes</td>
<td>Modification of target site through methylation.</td>
<td>Active efflux of antibiotic.</td>
<td>62</td>
</tr>
<tr>
<td>Aminoglycosides</td>
<td>Bacterial ribosomes</td>
<td>Modification of antibiotic through acetylation, phosphorylation, or adenylation.</td>
<td>Decreased uptake or increased efflux of antibiotic.</td>
<td>63</td>
</tr>
<tr>
<td>Oxazolidinones</td>
<td>Bacterial ribosomes</td>
<td>Mutations in the 23S rRNA of the ribosome, reducing the binding affinity of antibiotic.</td>
<td>Mutations in target enzymes</td>
<td>64</td>
</tr>
<tr>
<td>Fluoroquinolones</td>
<td>DNA gyrase and topoisomerase IV</td>
<td>Inhibits DNA synthesis and replication by interfering with DNA gyrase and topoisomerase IV.</td>
<td>Active efflux of antibiotic from the cell.</td>
<td>65</td>
</tr>
</tbody>
</table>
laboratories. Their mechanisms of action vary depending on the specific antimicrobial and the type of microorganism it targets. For example, bactericidal antimicrobials, such as penicillin, cephalosporins and vancomycin directly kill the bacteria. However, bacteriostatic antimicrobials on the other hand, such as chloramphenicol, tetracyclines and erythromycin, inhibit the growth of bacterial cells.

**AMR**

The initial indication of AMR became evident shortly following the identification of penicillin. In 1940, Abraham and Chain observed that a strain of *E. coli* had the capacity to render penicillin ineffective through the production of penicillinase. This early discovery laid the foundation for our understanding of resistance mechanisms. The timeline of antibiotic resistance development for different drugs are mentioned in Figure 1. When penicillin and sulphonamide were introduced in medical treatments during the late 30s and 40s of the 20th centuries, individuals believed that antibiotic use prevented every disease that is caused by the disease-causing agents. But the exploitation of these antibiotics has caused tremendous pressure in the development and improvement of AMR. Different bacteria started to develop resistance mechanism against the antibiotics and a few bacteria along with their mechanism of actions are mentioned in Table 1. The momentum for improving antibiotic resistance has been growing, especially as the use of antibiotics became more restricted after vancomycin was reserved for special cases, preventing resistance from developing. However, irregular antibiotic usage brought about the rise of methicillin-resistant *Staphylococcus aureus* (MRSA), which shows multi antimicrobial resistance. Consequently, vancomycin-resistant Enterococci (VRE) showed up in 1986 because of the far-reaching utilization of vancomycin, which was viewed as the antimicrobial after all other options had run out. AMR is primarily caused by the overuse and misuse of antimicrobial drugs. Over-prescription of antibiotics for viral infections, self-medication and over-the-counter sales of antibiotics contribute to the problem. In agriculture, antibiotics are not only used to treat sick animals but are also commonly added to the feed and drinking water of healthy animals to prevent illness. There is substantial evidence indicating that the use of antibiotics in animals significantly contributes to the development of AMR in human pathogens. Additionally, the slow development of new antimicrobial drugs, coupled with poor infection control practices in healthcare settings and communities, allows drug resistant pathogens to thrive and spread. These factors collectively contribute to the emergence and proliferation of AMR, making it a pressing global health concern. To battle these hazardous issues, it is essential for the continual efforts to develop new antimicrobial agents. Some recent antibiotics have been proven to be powerful upon MRSA and VRE, but these dangerous microorganisms will eventually develop resistance to these compounds.

**AMR mechanisms**

The resistance to antimicrobials is either inherited or intrinsic. For instance, in the natural resistance mechanism caused by *Pseudomonas aeruginosa*, the lower membrane permeability would be the probable reason for its intrinsic protection from numerous antimicrobials. Inherited resistance happens by securing a transposon or a plasmid that carries the genes that encode resistance mechanisms from other organisms, or via the process of chromosomal transformation.

The significant mechanisms of actions involved in the development of microbial resistance are:

- Reduction in drug binding affinity by altering drug targets.
- Modification of antibiotics causing inactivation of drugs.
- Increasing efflux or restricting uptake.
- Genetic mutations.

**Modification of drug targets**

The modification of drug targets is a crucial mechanism that microorganisms employ to develop AMR. This mechanism involves altering certain segments or structures in the microorganism that are the targets of the antimicrobial agents. For example, the change in...
the number and structure of penicillin-binding proteins (PBPs) is a major component of protection that is utilized by gram-negative microbes.  

PBPs are enzymes found in bacterial cell walls that play a key role in the synthesis and maintenance of the cell wall structure. Antibiotics like penicillin exert their antimicrobial effects by binding to these PBPs, thereby inhibiting cell wall synthesis and leading to bacterial cell death. However, when there is an increase in the production of PBPs or alterations in their structure, the binding of penicillin to these proteins may be hindered, reducing the drug’s effectiveness.

Another example of drug target modification seen in gram-negative bacteria, where resistance to vancomycin occurs due to changes in the structure of peptidoglycan precursors. Other mechanisms that have been reported include alterations in the surface charge of the bacterial cell membrane, such as a shift towards a positive charge which can interfere with the binding of calcium ions, thereby reducing the efficacy of certain drugs such as daptomycin.

**Inactivation of drugs**

The two fundamental manners by which microorganisms stop the activity of drugs are by either transferring a chemical group to the drug or by directly degrading the drug. One of the most well-known examples of drug inactivation is the production of β-lactamase enzymes by many bacteria. β-lactamases are an exceptionally enormous category of drug hydrolysing enzymes. These enzymes target and inactivate β-lactam antibiotics, which include widely used drugs like penicillin and cephalosporins. β-lactam antibiotics work by interfering with the synthesis of the bacterial cell wall, ultimately causing the cell to burst. However, β-lactamase enzymes can break the β-lactam ring, a key structural component of these antibiotics, rendering them inactive. As a result, the antibiotic can no longer disrupt the cell wall synthesis and the bacterium remains unharmed.

Tetracycline is another drug that can be easily hydrolysed and inactivated by the tetX enzyme which modifies the tetracycline molecule by adding a functional group to specific sites on the tetracycline structure. This change in structure interferes with the tetracycline’s ability to bind to the ribosome effectively. As tetracycline antibiotics depend on their specific interaction with the ribosomes to inhibit protein synthesis, the altered tetracycline molecule can no longer bind to the ribosome with the same affinity or effectiveness. As a result, it cannot disrupt the

**Figure 2. Different families of efflux pumps found in bacteria**
translation process and bacterial protein synthesis proceeds unimpeded.

The inactivation of certain drugs is brought about by a group of transferases responsible for transferring chemical groups such as phosphoryl, adeny1 and acetyl to these drugs.9 The most utilized mechanism is acetylation, which shows its effect against drugs like streptogramins, aminoglycosides and fluoroquinolones.

Active efflux

Many microorganisms have evolved with an amazing mechanism to counter increasing drug concentration levels in their cells.22 They simply efflux the antibacterial agents outside the cells through certain pumps in their cell surface known as efflux pumps.23 Via these molecular pumps, they can actively transport a wide variety of antimicrobial compounds and toxins out of the cells.24 Efflux pumps can exhibit specificity towards a single substrate or have the ability to transport a variety of structurally dissimilar compounds. This includes antibiotics from various classes, and these pumps may be linked to the phenomenon of multiple drug resistance (MDR).25

The efflux pumps come under five families which are:
- ATP-Binding Cassette (ABC) superfamily
- Multidrug and Toxic Compound Extrusion (MATE) superfamily
- Small Multidrug Resistance (SMR) superfamily
- Major Facilitator Superfamily (MFS)
- Resistance Nodulation and Cell Division (RND) superfamily

One of the major differences between the different families is the varying energy sources and assembly of the components as outlined in Figure 2. The ABC family relies on ATP hydrolysis for energy, as opposed to other transporters like MATE, MFS, RND and SMR superfamilies, which harness the proton-motive force from H+ or the electrochemical gradient of Na+ for energy to expel various compounds.26 Another difference as seen in the RND family is the tripartite complex comprising of the outer-membrane canal protein (OMP), an inner-membrane transporter and a membrane fusion protein (MFP) that bridges the OMP and inner-membrane transporter.26

The genes are chromosomally encoded and some of these are constitutively expressed while others are overexpressed under certain conditions like when the availability of a substrate is reasonable.27

Genes associated with efflux pumps can be obtained via intrinsic or acquired means. Certain bacteria possess these genes on their
chromosomes, providing an inherent survival mechanism in challenging environments.\textsuperscript{24} In contrast, other bacteria can procure these genes through diverse mechanisms such as mutations within local repressor genes, activation of a regulon controlled by a global transcriptional regulator, or the presence of efflux pump genes on plasmids.\textsuperscript{28} For example, the RND family efflux pumps play a crucial role in the intrinsic antibiotic resistance of gram-negative bacteria by actively expelling a diverse array of antibiotics and antimicrobials.\textsuperscript{29}

In gram-positive bacteria, the efflux pumps provide intrinsic resistance as they are encoded on the chromosome.\textsuperscript{23} Chromosomal encoding is responsible for MDR efflux pumps, exemplified by NorA, NorB, MepA and MdeA in \textit{S. aureus}.\textsuperscript{30} These pumps confer inherent antibiotic resistance in bacteria across a diverse spectrum. Conversely, certain efflux pumps in gram-positive are also carried around on plasmids or transposons, like QacA/B in \textit{S. aureus} or MefA and MefB in \textit{Streptococcus} spp., respectively.\textsuperscript{30}

**Genetic mutations**

In specific species, mutation is the primary, or sole, reason for AMR. Perhaps the best example would be Mycobacterium tuberculosis, where mutation is the primary cause of resistance to all clinical drugs in this bacterium. Combination treatment is required to treat tuberculosis which is a pathogen that causes prolonged sickness in humans.\textsuperscript{31} The risk caused by the mutation at the point of infection in microorganisms cannot be eliminated by the combination treatment which only diminishes the impact.

Alteration in their genome will prevent the ability of antibiotics to act on them thus creating a superbug or super bacteria. These naturally genetically modified bacteria produce resistance against the exploited drugs which then spread among their population leading to the entire species becoming resistant to certain drugs.\textsuperscript{19} Moreover, in some cases, after a bacterium dies, they may leave their genetic material outside which in turn might be picked
up by bacteria of a different species that might incorporate the usable resistant genetic material inside it resulting in producing new resistance.

**Superbugs**

Superbugs signify strains of MDR bacteria that are impervious to a large portion of the antibiotics and different drugs usually used to treat the disease they cause. The expression “superbug” was coined by the media and essentially refers to bacteria that has developed resistance to multiple drugs that were once effective against the infection it caused. Different methods of resistance mechanism developed by bacteria are mentioned in Figure 3. A couple of instances of superbugs include MRSA, VRE, multidrug-resistant _Pseudomonas aeruginosa_ and multidrug-resistant _Acinetobacter_. It is crucial to understand that any bacteria have the potential to evolve into a superbug and develop resistance to nearly any drug if those drugs are overused or misused. This misuse can have harmful consequences for crops, humans and animals. Thus leading to a situation where a multi-step approach may be the only effective approach to further prevent the spread and infection of these superbugs.

**Development of superbugs**

The development of superbugs is primarily due to the misuse and overuse of antibiotics, which leads to the emergence of drug resistant strains. When antibiotics are overused or misused, the microorganisms that cause the infections tend to develop resistance to multiple antibiotics via the various mechanisms that have been discussed earlier in this article. As the microorganisms evolve to survive the drugs that were previously lethal, they create antibiotic resistant strains and commonly used antibiotics become ineffective over time.

The emergence and spread of superbugs result from a combination of factors such as the overuse and misuse of antibiotics, inappropriate antibiotic prescribing practices and incomplete treatment courses. Nowadays the most common form of antibiotic misuse is consumption of

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**Table 2. Antibiotic Efflux Pump Families, their Substrates and Inhibitors**

<table>
<thead>
<tr>
<th>Efflux Pump Family</th>
<th>Substrates</th>
<th>Efflux Pump Inhibitors</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>ATP-Binding Cassette (ABC) superfamily</td>
<td>Tetracyclines</td>
<td>Reserpine</td>
<td>66, 67</td>
</tr>
<tr>
<td></td>
<td>Fluoroquinolones</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Macrolides</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Rifampicin</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Chloramphenicol</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Multidrug and Toxic Compound Extrusion (MATE) superfamily</td>
<td>Tetracyclines</td>
<td>Verapamil</td>
<td>67, 68</td>
</tr>
<tr>
<td></td>
<td>Fluoroquinolones</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Tigecycline</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Pentamidine</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Small Multidrug Resistance (SMR) superfamily</td>
<td>Tetracyclines</td>
<td></td>
<td>66, 67</td>
</tr>
<tr>
<td></td>
<td>Erythromycin</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Sulfadiazine</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Major Facilitator Superfamily (MFS)</td>
<td>Tetracyclines</td>
<td>Phenothiazines</td>
<td>66, 67</td>
</tr>
<tr>
<td></td>
<td>Fluoroquinolones</td>
<td>Thioridazines</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Erythromycin</td>
<td>Chalones</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Rifampicin</td>
<td>Piperine</td>
<td></td>
</tr>
<tr>
<td>Resistance Nodulation and Cell Division (RND) superfamily</td>
<td>Tetracyclines</td>
<td>Flavonoid’s</td>
<td>66, 67</td>
</tr>
<tr>
<td></td>
<td>Fluoroquinolones</td>
<td>Quercetin</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Erythromycin</td>
<td>Phenyl-arginine-β-</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Rifampicin</td>
<td>naphthylamide (PaβN)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>β-Lactams</td>
<td>Hydantoins</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Chloramphenico</td>
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</tbody>
</table>
antibiotics without consulting physicians. Furthermore, there are instances where healthcare professionals prescribe antibiotics unnecessarily. This may result from diagnostic uncertainty, patient requests, or precautionary practices. In cases where antibiotics are appropriately prescribed, some patients may not complete the prescribed course. Incomplete courses can lead to the survival of bacteria that develop resistance to the antibiotic. The reason for this lies in the fact that the most susceptible bacteria are eliminated first, leaving behind those that may possess greater resistance. Additionally, improper and excessive use of antimicrobials in livestock, fish farming and agriculture leads to the development of resistant bacteria, which causes major health risks.

Combating AMR

AMR may be a complicated issue with various causative factors. It’s the major reason behind health issues in the community, affecting people directly or indirectly. Given the necessity for widespread antibiotic use to combat AMR, the immediate development of new technology is imperative.

The overuse of certain antibiotics may result in the production of even more powerful superbugs which might ultimately become resistant to all known drugs. Hence new techniques to combat AMR are needed and to our surprise, many alternative techniques to combat AMRs have been developed by scientists all around the world to fight against the globally spreading AMR.

Targeting outer membrane

Bacteria have evolved various strategies to avoid contact with antibiotics, and one crucial mechanism involves preventing the entry of these drugs. Antibiotics typically exert their effects after entering the cell membrane, but bacteria have developed defenses to impede this process. The intricate structure of bacterial cell walls, comprising compounds like porins and lipopolysaccharides, serves as a barrier, hindering the entry of certain antibiotics. These compounds act as gatekeepers, preventing the intrusion of antibiotics. Few microbes undergo macromolecule mutations outside the cell membranes to prevent the entry of some antibiotics. However, certain antibiotics have been devised with means to overcome these barriers and enter the organisms.

Hydrophobic compounds

Certain compounds such as macrolides and rifampicin possess hydrophobic characteristics that allow the compounds to navigate through the cell membrane efficiently and easily cross the lipid bilayer.

Hydrophilic molecules

Antibiotics such as β-lactams, fluoroquinolones and phenicol antibiotics, are hydrophilic in nature. They leverage transport mechanisms by interacting with specific porins, allowing them to diffuse through the cell membrane and overcome the protective barriers.

Targeting β – Lactamases

Synthesis of enzymes that can counter the antibiotics administered is one of the major defense mechanisms employed by bacteria. β-lactam antibiotics which possess natural bactericidal properties were initially developed and widely used. However, the extensive use of β-lactam antibiotics led to the rapid development of resistant strains in gram-positive bacteria. The primary mechanism of β-lactam antibiotics involves the inactivation of transpeptidases, disrupting the final step in membrane biogenesis. Subsequently, bacteria developed mechanisms to evade β-lactam treatment such as reduced expression of transport proteins, increase of efflux pumps and expression of β-lactamases. As β-lactamases are enzymes that hydrolyse and disrupt β-lactams, this results in the inactivation of antibiotics that have the functional β-lactam group.

Several strategies can be employed to address this challenge. Firstly, developing inhibitors specifically designed to target and neutralize β-lactamases can help in preventing the degradation of β-lactam antibiotics. Additionally, a promising strategy is the use of combination therapy, wherein β-lactam antibiotics are administered alongside these newly developed inhibitors, enhancing the antibiotics’ effectiveness.
antibiotics represent another avenue, focusing on engineering changes that make these drugs less susceptible to β-lactamase activity. An example would be a β-lactamase-activated antibacterial prodrug where the prodrug only exhibits bactericidal properties upon activation by β-lactamase. By employing these methods, antibiotic resistance can be leveraged to specifically target bacteria that produce β-lactamase.

**Targeting efflux pumps**

The development of efflux pump inhibitors is one of the widely used strategies to combat resistance through efflux. These inhibitors are tiny molecules that attach themselves to the pumps and block the expulsion of the antibiotic compounds from the cell. An overview of several antibiotic efflux pump families, the specific antibodies associated with the efflux pumps and their inhibitors are mentioned in Table 2. As these small molecules do not possess intrinsic bactericidal activities, they have been tested for synergistic effects at various concentrations in combination with antibiotics.

Many ways have been explored for inhibiting efflux pumps, including:
- Disrupting channel protein assembly.
- Interfering with efflux pump gene expression.
- Preventing recognition by adding functional chains.
- Developing small molecules as substrates to block activity of efflux pumps.

Therefore, inhibition of efflux may result in several positive outcomes, including:
- Maintaining drug concentrations at therapeutic doses.
- Reducing multi-drug resistance.
- Reducing treatment periods.
- Enhancing the activity of antibiotics susceptible to efflux.

**Novel approaches to combat AMR**

Antibiotic resistance poses a significant threat to global human health. Imperative measures are needed to prevent the emergence and spread of multi-drug resistance organisms which could have far-reaching consequences. Novel methods to identify, validate and develop new techniques are necessary to overcome this serious issue. The latest breakthrough in addressing AMR is using genetic engineering tools. These tools, such as CRISPR/Cas9 technology and nanotechnology, can be used to directly target the antibiotic resistance genes of microorganisms. Figure 4 illustrates various delivery mechanisms for these techniques, which are currently at the forefront of efforts to combat antibiotic resistance.

**CRISPR/Cas9 to combat AMR**

CRISPR-Cas systems are known as bacterial adaptive systems and are analogues of the immune system. The CRISPR/Cas9 system has been employed in many fields of study, ranging from genome editing for agricultural purposes to disease eradication. Originally a natural defense mechanism employed by the bacteria to fight invading phage viruses, scientists have harnessed and replicated these techniques to develop a technology aimed at preventing bacterial impact on humans. Bacterial genes contain CRISPR/Cas arrays, which consist of repeating sequences of DNA, known as repeats which are separated by unique sequences of equal sizes known as spacers. In the context of combating AMR, CRISPR/Cas9 technique is specifically programmed to target the DNA at specific sites, consequently cleaving AMR genes resulting in the disruption of the bacterial cells’ AMR mechanism.

**Nanoparticles as a tool to combat AMR**

Nanotechnology is an emerging field that utilizes nanoparticles to perform multiple or specific functions. These nanoparticles can be customized to perform specific tasks. In the context of AMR, utilizing nanoparticles offers a promising approach and unique advantages to address challenges faced by traditional antimicrobials. Nanomaterials have the capability to interfere with bacterial membranes, target intracellular components, and facilitate the delivery of antimicrobial agents. This results in improved therapeutic effectiveness against stubborn MDR infections. The use of nanotechnology alongside CRISPR/Cas9 holds promise for advancing treatments against MDR bacterial infections. As mentioned earlier, CRISPR/Cas9, known for its precise targeting, can disrupt drug resistance genes used by microbes for infection or directly eliminate pathogens. However, despite its efficacy, challenges in
in-vivo delivery efficiency have limited its broader application. Nanotechnology provides a solution by enhancing the effectiveness and safety of CRISPR/Cas9 components through specialized nanoparticle delivery systems, addressing these delivery shortcomings.58

CONCLUSION

AMR poses a severe global threat that could lead to a catastrophe if not promptly addressed. The most effective strategy against disease caused by bacterial pathogens is not by developing antibiotics to eliminate the pathogens but also to employ necessary measures to control the use of antibiotics. It is crucial for individuals worldwide to contribute to the fight against AMR by preventing the misuse of antibiotics in human and animal treatment. Since this cannot be accomplished overnight, this article offers several unique and successful approaches to combat MDR bacteria. Along with the development of new techniques to fight against AMR may incur significant costs that may be prohibitive for ordinary individuals, the emphasis should be on regulating antibiotic usage. Addressing the crisis of AMR involves raising public awareness on antibiotic misuse and overuse. Accessible diagnostic tools play a crucial role in ensuring that the proper antibiotics are prescribed to the right patient at the correct time, in regulated dosages for a particular time frame until their immune system effectively responds to the infection. Implementing antibiotic stewardship programs and regulatory measures are essential to control antibiotic use in both medical and agricultural settings. These programs can aid in ensuring that there is no room for bacteria to develop resistance mechanisms, ultimately preventing a widespread crisis.

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CONFLICT OF INTEREST

The authors declare that there is no conflict of interest.

AUTHORS’ CONTRIBUTION

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DATA AVAILABILITY

All datasets generated or analyzed during this study are included in the manuscript.

ETHICS STATEMENT

Not applicable.

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