Antibiotic Resistance Creating New Epoch

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Abstract

Antibiotics are an efficient means of treating bacterial infections. They help in the reduction of mortality and morbidity. Discovery of antibiotic has created a new epoch where infectious diseases have been put to an end. But the organisms have developed resistance during the course of evolution due to mutations. The major problem which is being faced is the treating of infectious diseases caused by microbes which have developed resistance. The antibiotic resistance is being carried by the organisms due to acquisition of resistance genes to next generations by horizontal gene transfer. The present review is focused on what are the various mechanisms adopted by the organisms against antibiotics and new techniques involved in treating the infectious diseases and methods to overcome the antibiotic resistance.

Keywords: Antibiotic resistance, Multi Drug Resistance [MDR]; Methicillin Resistance Staphylococcus aureus [MRSA]; XDR-TB

Introduction

Antibiotics are the chemical substances that are antagonistic to the growth of microorganisms which are ubiquitous in nature. They are either produced by microorganisms or prepared synthetically. Penicillin was the first antibiotic discovered by Alexander Fleming from the fungi named Penicillium [1]. All penicillins are β-lactam antibiotics (Figure 1) used to treat bacterial infections in clinical medicine. Cephalosporins & Carbapenems are also included in β-lactam antibiotics. They are classified into two groups based on their biological activity on microorganisms Bactericidal which kills the microbes, Bacteriostatic (erythromycin) [2] which inhibits the growth of microorganisms.

Antibiotics are classified based on spectrum.

• Wide / Broad spectrum
• Narrow spectrum

Broad spectrum antibiotics effects several types of bacteria and fungi, they acts both on Gram-negative and Gram-positive Bacteria, where as Narrow spectrum antibiotics are specific towards some bacterial families. Penicillins fall under broad spectrum of antibiotics which are effective against a wide range of bacteria. And hence the consistent use of such antibiotics has led to the emergence of resistance of antibiotics in microbes creating new epoch.

Antibiotics include Antibacterials, Antifungals, Antimalarials, Antivirals.

Antibacterials

The compounds that act against the bacteria are called antibacterials. Antibacterials can kill or inhibit the growth of bacteria. Antibacterials are the most important medications used in health care [3]. They play a pivotal role in combating many bacterial infections. Most of the Antibacterials are semisynthetic natural compounds like penicillins, carbapenems, cephalosporins. Whereas sulfonamides, Quinolones and oxazolidinones are purely synthetic compounds. Ofloxacin [OFX] is antimicrobial agent with a broad antimicrobial spectrum against Gram-positive and Gram-negative bacteria and is considered safe. It interferes with the enzyme DNA gyrase which is essential for synthesis of DNA [4]. Silver nanoparticles are emerging trends in nanosciences technology which are used as antibacterials to fight infectious organisms [5]. The Potential pathogenic bacteria are administered into the body by ingestion or through eyes when rubbed with contaminated fingers [6]. And this type of administration of pathogens can be restricted by using disinfectants.

Antifungals

The compounds which are antagonistic to the growth of fungi are called Antifungals. Fungal organisms can cause great harm and damage. They infect people, animals and plants, producing diseases that range in seriousness from mild infection to death [7]. Mycosis is a fungal infection caused in animals and humans. Inhalation of fungal spores or localized colonization of the skin may initiate persistent infections, therefore mycoses often start in the lungs or on the skin. Invasive mycoses associated with high morbidity and mortality rates are increasing among immunocompromised or severely ill patients [8].

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Various types of antifungals are in use to treat fungal infections and diseases.

Polyene antifungals: A molecule with multiple conjugated double bonds is called polyene. This interacts with sterols of cell membrane of fungi which makes channels in the membrane ejecting the monovalent ions and small organic molecules out of the cell leading to the death of the cell.

Azole antifungal drugs: Fluconazole, itraconazole, and ketoconazole inhibit cytochrome P450-dependent enzymes (particularly C-14-demethylase) involved in the biosynthesis of ergosterol, which is essential for fungal cell membrane structure and function. Voriconazole is a fluconazole with enhanced antifungal spectrum. It is like other azole antifungals, alters the cell membrane function and permeability, resulting in cell dysfunction [9]. Voriconazole is a safe promising and cost effective antifungal [10].

Anti malarials

Compounds which act against malaria causing Plasmodium falciparum are called antimalarials [11]. Malaria is caused by a parasite from plasmodium species that is passed from one human to another by the bite of infected Anopholes mosquitoes. Malaria involves high fevers, shaking chills, flu-like symptoms, and anemia. The parasites after infection are called sporozoites travel through blood stream to liver and mature over there and release merozoites. The parasites multiply in the red blood cells.

The major symptoms observed in malaria are:

- Merozoites are released into blood stream
- Hemoglobin is released into circulation on breakage of red blood cells
- Anemia as a result of destruction of red blood cells

Hydroxychloroquine & chloroquine are anti-malarial medications used effectively active against malaria. Quinine is an alkoid which acts as a blood schizonticidal. Quinine is less effective and more toxic as a blood schizocidial agent than chloroquine.

Due to the acquired global resistance of malaria parasites to antimalarial drugs, there is need for new approaches to malaria treatments. Targeting of enzymes which play pivotal roles in the parasite’s life cycle is being executed to overcome the resistance of the organism [12]. Treatment with cysteine protease inhibitors holds hemoglobin hydrolysis and parasite development in vitro.

Antivirals

The compounds which are antagonistic to the viral particles are called antivirals. Viruses are constantly plaguing the world, causing severe infections and mortality [13]. Dengue virus is transmitted to humans through bites of mosquito species, Aedes aegypti and A. albopictus [14]. The emergence of antiviral drugs is the expanded knowledge of the genetic and molecular function of organisms by biomedical researchers. Sulfated polysaccharides (sPS), in particular dextran sulfate (DS) and related polyanionic compounds have been shown to be active against various viruses [15].

Mechanism of action of Antiviral:

- In Influenza A virus uncoating is inhibited by amantadine and rimantadine
- Gamma globulins neutralize the viral particles
- Viral DNA polymerase is inhibited by guanosine derivative Acyclovir in Herpes virus.
- Gancyclovir is a guanosine derivative which is phosphorylated and incorporated into viral DNA inorder to suppress its replication in cytomegalovirus.
- Neuramidase is an enzyme essential for virus replication; this is inhibited by the drugs Zanamivir and Oseltamivir in H1N1 & H5N1 cases.
- Interferon’s and Monoclonal antibodies are found active against viral infections.

HAART is an antiretroviral therapy in use nowadays. It consists of combination of 3 antiviral drugs with one protease inhibitor [16]. Since this therapy is in use the decrease in mortality and morbidity is observed [17].

Disadvantages of antibiotics

- They cause allergic response inducing hyper sensitivity.
- The possible side effects are observed in the treatment of diseases, although these should be rare at the doses and length of time of treatment [18].
- Antibiotics destroy friendly bacteria present in the body that helps in detoxification, elimination of waste and cleansing of the blood and Liver.
- The persistent use of antibiotic to the bacteria causes bacteria accustomed with it and no longer get effected by antibiotics, transforming into drug resistant bacteria.
- The bacterium acquires the resistance gene and transfers them to next generations, multiplying the drug resistance bacteria.
- Antagonism between antibiotics contributes to the disadvantages of the inappropriate use of antimicrobial combinations [19].

Antibiotic Resistance

Antimicrobial resistance is the resistance of microorganisms to antimicrobials like antibiotics and drugs. Infectious diseases caused by microorganisms which are resistant to antimicrobials often results in prolonged illness and greater risk of death. Antibiotic resistance bacteria have posed severes threat to public health [20].

Organisms that display multidrug resistance are

- Pathologic cells.
- Neoplastic cells.

Pathologic cells include bacterial cells

MDR & XDR-TB are the two deadly forms of TB which are resistant to the drugs used in treating TB and imposing threat to human beings [21,22]. Streptococcus pneumonia is an important pathogen causing a variety of diseases including pneumonia, meningitis, and otitis [23]. Staphylococcus aureus is a major cause of various hospital acquired infections. These infections range from superficial skin infections to deeper infections of hair follicles, abscesses, and deep tissue infections and even to systemic infections including those of the heart, lungs, bones, and blood [24]. Staphylococcus aureus is an important pathogenic facultative Gram-positive bacterium [25] which has acquired resistance; it also forms biofilms which are the bacterial community that encloses themselves in a polymer matrix [26]. The adeB gene found in MDR strains confers resistance to aminoglycosides and tetracyclines in Acinetobacter baumannii [27]. Pseudomonas aeruginosa is an emerging opportunistic and nosocomial pathogen infecting only compromised
tissues and causing pathology in the gastrointestinal tract [28]. Drug resistant mycobacterium is increasingly common in different parts of the world and is fueled with spread of Acquired Immunodeficiency Syndrome (AIDS) [29].

Eg. Meficillin Resistant Staphylococcus aureus (MRSA), Pseudomonas aeroginosa, Acinetobacter baumannii, Mycobacterium tuberculosis.

Neoplastic cells include tumor cells

Multidrug resistance, the principal mechanism by which many cancers develop resistance to chemotherapy drugs, is a major factor in the failure of many forms of chemotherapy. It affects patients with a variety of blood cancers and solid tumors, including breast, ovarian, lung, and lower gastrointestinal tract cancers. Tumors usually consist of mixed populations of malignant cells, some of which are drug-sensitive while others are drug-resistant. Chemotherapy kills drug-sensitive cells, but leaves behind a higher proportion of drug-resistant cells. As the tumor begins to grow again, chemotherapy may fail because the remaining tumor cells are now resistant.

Causes of antibiotic resistance

• Antibiotic abuse
• Horizontal Gene Transfer

Antibiotic abuse: Drug abuse has become increasingly common these days [30]. Inappropriate use of prescriptions for antibiotics creates new, drug-resistant strains of common diseases. Prescribing antibiotics carefully and using them wisely is the key to preventing the spread of antibiotic-resistant illnesses. The premature breaking of the antibiotic doses by patients is also a major cause of antibiotic abuse.

The Antibiotics were given along with the feed to the livestock in order to reduce the infections and increase the rapid growth of the animals. This ultimately increases the risk of germs which pass onto humans making them resistant to antibiotics. Implementation of reduced prescription of antibiotics showed a remarkable decrease in drug resistant strains [31].

Horizontal gene transfer: A spontaneous genetic mutation confers resistance to the drugs in bacteria. Generally more than one mutation is required to provoke drug resistance. Many of the antibiotic resistant genes reside on plasmids which are passed onto next generations through horizontal gene transfer (Figure 2). The resistant genes are conceded to the next generations by transformation, transduction or conjugation, thus multiplying the population of resistant bacteria.

Drug resistance in Neoplastic cells: Chemotherapy is used to treat cancer. Some of the cancer cells which were not killed by chemotherapy mutate and the mutated cells proliferate passing on the drug resistance to the newly formed cells. Apoptosis and Senescence are the two main programs which help the drugs in treating cancer. Mutations which alter these programmes will lead to chemoresistance in tumor cells.

Cancer is the continuous proliferation of the cells in the body. The lumps or masses produced due to uncontrolled growth are called tumors [32]. Cancer is of two types Benign and Malignant. Uncontrolled invasive growth is main characteristic of malignancy, and metastasis is in most cases the reason why cancer patients die [33].

Tumor cells contain oncogene that has ability to cause cancer. Apoptosis is the programmed cell death practiced by many normal cells. Resistance to apoptosis leads to the progression of solid cancers [34]. Numbers of factors are responsible for cancer genesis. Loss of control of cell cycle, Angiogenesis, Cellular neoplastic transformation, Resistance to apoptosis, and acquisition of invasive properties initiate the growth of solid tumors [35]. Chemotherapy is used to treat cancer. Chemotherapy means use of chemical compounds in the treatment. The persistent use of chemical compounds has resulted in resistance of cancer cells. Resistance could also result from alterations in cell proliferation [36].

The major Hindrance in cancer treatment is acquired resistance to chemotherapy, there are several methods employed by cancer cells to resist the treatment.

• Cells which are not killed by chemotherapy mutate and develop drug resistance.
• Over production of proteins by gene amplification renders anticancer drug ineffective
• Cancer cells drive out the drug before it reaches the targeted site
• Transport channels across the membrane stop working and inhibits the drug uptake
• Patients with circulating tumor cells expressed MRP1 and MRP2, two drug-export pumps responsible for anthracyclines efflux[37].

Various other means of treatment are being investigated to fight cancer. Many new techniques has evolved to treat cancer. Neoadjuvant Chemotherapy is in practice to treat breast cancer [38]. Neoadjuvant chemotherapy is done prior to surgery making the cancerous cells shrink in size giving a clear distinction between cancer cells and healthy tissue surrounding it in order to decrease the damage caused to healthy tissue during surgery [39].

Bacterial cells interact with tumor cell lines in brain tumor which advances the treatment of brain tumor with bacterial cells [40]. γ-Tocotrienol is one of the eight natural isoforms that make up the family of vitamin E compounds and displays potent antiproliferative and apoptotic activity against neoplastic mammary epithelial cells [41]. Coordination compounds have found to be effective as antineoplastic compounds[42]. Agaricus sylvaticus assess the antioxidant potential which helps in preventing pre mature aging and cancer [43]. Angio-inhibitors are also available naturally. Consumption of natural angi-inhibitors can help prevent cancer [44]. miRNAs can delineate cervical cancer from normal cervical tissue, and miRNAs have potential as markers for progression from dysplasia to invasive cervical disease. EGF, PDGF as well as their receptors are implicated in experimental and human malignant diseases. Wounds and developing tumors are biologically similar niches of dynamic interaction between varieties of cell types sharing many histological features. For tissue repair and tumorigenesis, cell proliferation, migration, survival, and angiogenesis are
instrumental events whereas all these processed appeared governed by a plethora of growth factors.

**Acquisition of resistance**

Microbes play different strategies to develop resistance some of them include.

**Enzymatic Inactivation**: β-lactam is a four membered ring. It is a part of core structure of many broad spectrum antibiotics. The enzyme β-lactamases [45] hydrolyses the β-lactam ring disturbing the molecular structure of the antibiotics. Carbapenems are the antibiotics which are very efficient against the Gram-negative resistant bacteria. They are inhibited by the enzymes carbapenemases. Carbapenemases are the enzymes which confer resistant to carbapenem in Gram-negative Bacteria such as *Pseudomonas aeruginosa*. MBL are the metallo-beta-lactamases (IMP, VIM, SPM, GIM types) and are clinically significant carbapenemases. Carbapenemases are also active against ammoglycosides and quinolones. MBLs contain Zn\(^{2+}\) cation. They belong to the molecular class B family. They mediate resistance to beta lactams by cleaving the amide bond in beta lactam ring.

**Drug impermeability**: Organisms with high permeability can develop resistance by decreasing the permeability. In *Pseudomonas aeruginosa* resistance is achieved by lacking the channel OprD which allows β-lactamase to pass through it. The hospital isolates of *Pseudomonas aeruginosa* has developed resistance by using this mechanism. The permeability depends on physicochemical properties of the outer membrane of the bacteria. Hydrophobic antibiotics are effective against Gram-positive bacteria rather than Gram-negative Bacteria. The lipopolysaccharide layer contains less fluidity and confers rigidity to the outer membrane which doesn’t allow the hydrophobic antibiotics penetrate through it. In Gram-negative bacteria diffusion of Hydrophilic antibiotics is accomplished by diffusion channels formed with the help of porin proteins. Nutrients, Products of metabolism, and cephalosporins drugs are diffused through porin channels. Lack of these porins ie; OmpF and OmpC [46] confers resistance to cephalosporins.

**Efflux pumps**: Active efflux is a mechanism responsible for extrusion of toxic substances and antibiotics outside the cell, this plays a pivotal role in xenobiotic metabolism. This mechanism contributes to bacterial antibiotic resistance. Efflux systems function via an energy-dependent mechanism (Active transport) to pump out unwanted toxic substances through specific efflux pumps. Tetracycline efflux is achieved by an export protein from the major facilitator super family (MFS) [47]. The export protein was shown to function as an electroneutral antiport system which catalyzes the exchange of tetracycline-divalent metal-cation complex for a proton. The distinct energy dependent transporters called efflux pumps extrude the antibiotics from the periplasm to the exterior of the cell. This efflux pump overproduction is generally accompanied by an increase in resistance to two or more structurally unrelated antibiotics and contributes to the emergence of multidrug pathogens.

**Biofilms**: Biofilms are the important survival mechanisms of bacteria. Biofilms protect bacteria against antibiotics and phagocytosis [48]. The microbes adhere to the surface and colonise. They can anchor more efficiently and permanently with the help of pili. They are mostly found on submerged areas and substrates in contact with aqueous solution. They are mostly found on submerged areas and substrates in contact with aqueous solution. The RapA protein belongs to the SWi/SNF superfamily of helicase-like proteins, which has been implicated in chromatin remodeling in eukaryotic cells. In the RapA-deficient mutant, several genes were regulated, including the yhcQ and yeeZ genes. The yhcQ is putatively concerned with encoding a multidrug resistance pump and yeeZ with an unknown envelope function. The enhanced penicillin resistance of the wild type biofilm is due to a dual strategy: impaired penetration of the biofilm through its matrix, and rapid efflux of the antibiotic that still manages to penetrate. OprF is involved in adherence to eukaryotic Cells and in biofilm formation under anaerobic conditions [49].

**How to overcome the problem?**

- Reducing the disease burden and the spread of infection
- Improving access to appropriate antimicrobials
- Improving use of antimicrobials
- Strengthening health systems and their surveillance capabilities
- Enforcing regulations and legislation encouraging the development of appropriate new drugs and vaccines.
- One area that has potential for development of new drugs is the formation of metal coordination complexes [50].

To combat antibiotic resistance is to strengthen the action of existing antibiotics by modifying them so the bacterial enzymes that cause resistance cannot attack them. Alternately, “decoy” molecules can be used along with the antibiotic, so that the bacterium’s resistance enzyme attacks the decoy molecule rather than the antibiotic. Decoy molecules such as clavulanic acid or sulbactam are already in use for blocking the beta-lactamase enzymes that destroy the penicillin family of drugs.

An alternative approach to the antibiotic resistance problem is to interfere with the mechanisms that promote resistance, rather than to attempt to kill the bacteria. For example, interfering with the duplication or movement of a bacterium’s genetic material would eliminate the transfer of resistance genes between bacteria.

**Re-engineered Drugs (Vancomycin)**

The new vancomycin analogue can grab hold of the mutant peptidoglycan, and again prevent the bacteria from making the cell wall and killing the resistant bacteria. The redesigned antibiotic maintains its ability to bind the wild type peptidoglycan as well. Changing the properties of a key amide at the core of the natural products structure required a new synthetic strategy that only the most talented chemists could achieve in the lab. The preparation of the entire structure took a great deal of time and a fresh approach.

**Apitherapy**

The Medical use of Honey and Honey Bee products like bee bread, bee venom, propolis, royal jelly, apilarnil is called Apitherapy. Honey has a potent antibacterial activity and is very effective in clearing wounds, infection by inhibiting a wide range of bacteria [51]. The in vitro studies of honey revealed its antitumor and antifungal properties [52]. Melittin is the active component of bee venom which possess anti bacterial, antiinflammatory & antiviral properties. Bee venom consists of peptides and proteins which have neurotoxic and immunogenic effects. Apitherapy has become the focus of interest as a form of alternative and preventive medicine for treatment of a number of clinical cases. Honey has been suggested as an effective healing agent in treatment of post-operative wounds. In many cases honey has cleaned up wounds when conventional treatment was unsuccessful. Commercial honey has been reported to accelerate wound healing when applied topically to experimentally induced wounds infection. Antibacterial activity of honey was attributed to hydrogen peroxide naturally present in it. The antiproiferative activity of Tualang honey was observed in treatment of Oral squamous cell carcinomas (OSCC) and Human Osteosarcoma (HOS).
Surgery and Radiotherapy of OSCC & HOS leads to reduced quality of life. Apoptotic cellular changes like becoming rounded, reduction in cell number, blebbed membrane and apoptotic nuclear changes like nuclear shrinkage, chromatin condensation and fragmented nucleus are observed in OSCC and HOS cell lines [53].

**Phage therapy**

Viruses carry only a minimal number of genes and thus, for replication they must pirate the cellular machinery of infected cells [54]. Bacteriophage is a virus that depends on bacterial cellular machinery for its survival [55]. The potential use of the phage for the control of both local and systemic human S. aureus infections has been observed in recent studies [56]. Bacteriophages are harmless not only to the host organism, but also to other beneficial bacteria, such as gut flora, reducing the chances of opportunistic infections. Bacterial phage therapy has proved extremely successful in the treatment of topical infections, as has the inhalation of phages for the treatment of lung infections. The ability of phages to prevent biofilm formation on medical devices has received much attention, mainly in the area of catheter coatings [57]. Phage therapy is considered to be safe compared to use of antibiotics. The bacteria release endotoxins when destroyed within the patients, this causes fever and toxic shock in extreme conditions. To overcome this situation genetically engineered bacteriophages are used which lacks endolysin producing gene, this keeps the destroyed bacteria intact because lysis stage is disabled. And the phages are phagocytosed by the phagocytes.

**Photodynamic therapy**

Light-activated antimicrobial agents (photosensitizers) are promising alternatives to antibiotics for the treatment of topical infections. To improve efficacy and avoid possible damage to host tissues, targeting of the photosensitizer to the infecting organism is desirable, and this has previously been achieved using antibodies and chemical modification of the agent. The study has investigated the possibility of using a bacteriophage to deliver the photosensitizer tin (IV) chlorin e6 (SnCe6) to *Staphylococcus aureus*. SnCe6 was covalently linked to *Salmonella* bacteriophage 75, and the ability of the conjugate to kill various strains of *S. aureus* when exposed to red light was determined. Substantial kills of methicillin- and vancomycin-intermediate strains of *S. aureus* were achieved using low concentrations of the conjugate (containing 1.5g/ml SnCe6) and low light doses (21J/cm2). Under these conditions, the viability of human epithelial cells (in the absence of bacteria) was largely unaffected. On a molar equivalent basis, the conjugate was a more effective bactericide than the unconjugated SnCe6, and killing was not growth phase dependent. The conjugate was effective against vancomycin-intermediate strains of *S. aureus* even after growth in vancomycin. The results demonstrated that a bacteriophage can be used to deliver a photo sensitizer to a target organism, resulting in enhanced and selective killing of the organism. Such attributes are desirable in an agent to be used in the photodynamic therapy of infectious diseases.

**Active Plant Components for Therapeutic Use (Phyto Therapy)**

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**References**


