Mini review on antimicrobial agents

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Abstract
Antimicrobial agent is a general term that is mainly concerned with antibiotics, antibacterials, antifungals, antivirals and antiprotozoans. The antimicrobial agent is a substance that kills or inhibits the growth of microorganism like bacteria, fungi and protozoan. Antimicrobial agents are capable of acting by two modes either by inhibiting the spread of the infectious agent (microbiostatic) or by killing it outright (microbicidal). The treatment of using antimicrobial agents is currently facing its own limitation, 1, 2 due to the development of resistance by the microbes over the period of time.

Keywords: drugs, activities, antifungal, antibacterial

Introduction
Fungi: The fungus is a eukaryotic organism which is classified as a separate kingdom from plants, animals and bacteria. The fungus kingdom is divided into one subkingdom, seven phyla, and ten subphyla having a vast diversity of taxa with varied ecologies, life cycle strategies, and morphologies ranging from single-celled aquatic chytrids to large mushrooms. They are omnipresent, comprising an estimated 250,000 species, of which only about 150 have reported to cause disease in humans. 3-5 Fungi contain unicellular, multinucleate, and multicellular forms and are classified on the basis of their reproductive spores and the nature of hyphae. They divide sexually or asexually or by both ways 1.2. Fungi are almost entirely multicellular with exception of yeast Saccharomyces Cerviseae which is prominent unicellular fungus. Fungi are heterotrophic, deriving their energy from another organism, whether alive or dead. Fungi are eukaryotic protista differing from bacteria in many ways, as they possè’s rigid cell wall containing chitin, mannan, other polysaccharide’s and their cytoplasmatic membrane contains sterol. The major difference between fungal cells and plant cell is that fungal cell walls contain chitin, while plants contain cellulose 3-5.

Mycology
The discipline of biology which is devoted to the study of fungi is known as mycology. Mycology is concerned with the systematic study of fungi, including their genetic and biochemical properties 1.3-7. Mycoses affecting humans can be divided into four groups based on the level of penetration into the body tissues as,

A. Superficial mycoses, caused by fungi growing only on the outermost surface of the skin or hair. An example of a fungal infection is Tinea versicolor, a fungus infection that commonly affects the skin of young people, especially the chest, back, upper arms and legs.

B. Cutaneous mycoses or dermatomycoses, caused by fungi growing only in the superficial layers of skin, nails, and hair causing infections commonly known as athlete's foot, jock itch, and ringworm.

C. Subcutaneous mycoses, caused by fungi penetrating below the skin in the subcutaneous, connective, and bone tissue. The most common is sporotrichosis, occurring amongst gardeners and farmers who come in direct contact with soil.

D. Systemic or deep mycoses, are caused by primary pathogenic and opportunistic fungal pathogens. The primary pathogenic fungi cause infection in a normal host; whereas, opportunistic pathogens require immune depressant host in order to establish infection (e.g., cancer, surgery and AIDS). The primary pathogens usually gain access to the host via the respiratory tract and include Coccidioides immitis, Histoplasma capsulatum, Blastomyces dermatitidis and Paracoccidioides brasiliensis. The opportunistic fungi invade via the respiratory and alimentary tract, and include Cryptococcus neoformans, Candida spp., Aspergillus spp., Penicillium marneffei, Zygomycetes, Trichosporon beigelii and Fusarium spp.

Mycology - pathogenic fungi
The study of pathogenic fungi is referred to as medical mycology. pathogenic fungi cause disease in humans or other organisms. The commonly observed pathogenic fungi are enlisted below 3, 5.

A. Candida: Candida species are important human pathogens causing opportunistic infections in immune compromised hosts (e.g. AIDS sufferers, cancer patients and transplant patients). The infections caused by the Candida species are difficult to treat and could be fatal. Candida species alone account to 30-40% death cases caused by the systemic infections. Fungal species of the genus Candida generally live communally on and in the human
body. There is increase in development of drug resistance by \textit{Candida} species to current therapies, motivating researchers to understand their genetics and discover new therapeutic targets.

B. \textit{Aspergillus}: The aerosolized \textit{Aspergillus} spores are found almost everywhere around human being and generally does not posses health issues. But still \textit{Aspergillus} is capable of causing disease in three major ways: by the production of mycotoxins; by the induction of allergic responses; and lastly by the localized or systemic infections. \textit{Aspergillus flavus} produces mycotoxins, aflatoxin which can act as both toxin and a carcinogen, having ability to contaminate foodstuff such as nuts.

C. \textit{Cryptococcus}: The majorities of \textit{Cryptococcus} species are found in the soil and generally do not cause disease in humans. Exception is \textit{Cryptococcus neoformans} which cause disease in immune depressant patients like AIDS, causing a severe form of meningitis and meningo-encephalitis.

D. \textit{Histoplasma}: \textit{Histoplasma capsulatum} can cause histoplasmosis in humans, dogs and cats. The infection is usually due to inhaling the contaminated air and is prevalent in USA and Asia.

E. \textit{Pneumocystis}: \textit{Pneumocystis jirovecii} can cause a form of pneumonia in people with weakened immune systems, such as the elderly, AIDS patients and premature born children.

F. \textit{Stachybotrys}: \textit{Stachybotrys chartarum} can cause respiratory damage and severe headaches, in houses that are persistently damp.

1.3.2 Fungal infection prominent in particular diseases
The association of most commonly occurring fungi along with the patient suffering from the particular disease is enlisted below,
- \textit{Zygomycetes}, \textit{Rhizopus}, \textit{Mucor}, \textit{Absidia}: Diabetes.
- \textit{Candida}, \textit{Cryptococcus}, \textit{Histoplasma}: AIDS.

1.3.3 Clinically significant fungi and the site they affect
The pathogenicity and virulence of fungi causing infections, in humans is major concern in clinical world which focuses on the major causative agents of disease, particularly \textit{Candida}, \textit{Cryptococcus} and \textit{Aspergillus} spp. The large diversity of potentially harmful fungi existing outside these groups, though rare may still posses’s potential to be more important than the common clinical fungi. The site affected by the clinically significant fungi are enlisted as below,
- \textit{Malassezia furfur} and \textit{Exophiala werneckii}: Superficial skin.
- \textit{Piedraia hortae} and \textit{Trichosporon beigeli} : Hair.
- \textit{Microsporum} species: Skin and hair.
- \textit{Epidermophyton} species: Skin and nails.
- \textit{Trichophyton} species: Skin, hair and nails.
- \textit{Histoplasma capsulatum}, \textit{Penicillium} species: Systemic respiratory.

- \textit{Blastomyces dermatitidis}: Subcutaneous/respiratory.
- \textit{Cryptococcus neoformans}: Respiratory/CNS.

Antifungal agents: The antifungal agents are classified according to their mode of interfering with
1. Cell wall synthesis,
2. Plasma membrane integrity,
3. Nucleic acid synthesis,
4. Ribosomal function.

The (Figure 1) gives us brief idea about the mode of action antifungal drugs [5-10].

Fig 1: Classification of antifungal agents by their mode of action.11

The classification of antifungal drugs are made upon their mode of action, viz.,

A. Systemic Antifungal Drugs
1. Polyenes antibiotics: Amphotericin B.
2. Azole derivatives
   - Imidazole: Ketoconazole, Miconazole.
   - Triazole: Fluconazole, Itraconazole, Voriconazole, Posaconazole, Ravuconazole.
3. Echinocandin: Capsofungin, Anidulafungin, Micafungin.
5. Nikkomycin

B. Topical Antifungal Drugs
3. Others: Tolnaftate, Undecyclenic acid, Povidone iodine, Triacetin, Gentian violet, Sodium thiosulphate, Cicloporeoxalamine, Benzic acid, Quinidochlor.

Systemic Antifungal Drugs for superficial infections

Class of drug with general mechanism of action: The general mechanism of action of the drug depending upon its class is shown in (Figure 2) and the drugs are as enlisted below [1, 4, 5],

1. Griseofulvin: The griseofulvin was the earliest known inhibitory agent specific to fungal species. The griseofulvin binds to tubulin, interfering with the functioning of
microtubule and thus inhibiting mitosis. It binds to keratin in keratin precursor cells and makes them resistant to fungal infections. It is only when hair or skin is replaced by the keratin-griseofulvin complex that the drug reaches its site of action.

2. **Flucytosine:** Flucytosine works as an antifungal agent by the conversion to 5-fluorouracil within target cells by cytosine deaminase, thereby inhibiting synthesis of DNA through effects on thymidylate synthase in fungi. For this mechanism of action, the target cells must possess certain enzymes like cytosine permease to internalise the flucytosine molecule, cytosine deaminase to convert it to 5-fluorouracil and uracil phosphoribosyl transferase to convert 5-fluorouracil into a substrate for nucleic acid synthesis. The use of flucytosine is restricted to pathogenic yeasts (Candida species and Candida neoformans) as majority of the filamentous fungi lack these enzymes.

3. **Polyenes:** The mode of action of polyene is different than other antifungal agents, instead of inhibiting an enzyme, it binds to ergosterol, which is the principal sterol in fungal membranes, thereby perturbing membrane function to the point of causing leakage of cellular contents. As with other polyene antifungals, amphotericin B associates with ergosterol, forming a transmembrane channel that leads to monovalent ion (K⁺, Na⁺, H⁺ and Cl⁻) leakage, which leads to fungal cell death.

4. **Azoles:** The largest and most widely used class of antifungal agents is that of the azole-antifungals. Azoles inhibit 14-α demethylation of lanosterol in ergosterol biosynthetic pathway. This inhibition prevents the conversion of lanosterol to ergosterol, which is an essential component of the fungal cytoplasmic membrane and subsequent accumulation of 14α-methyl sterols.

5. **Allylamines:** At this moment there are two allyamines naftifine and terbinafine, and one thiocarbamate antifungal, tolnaftate which are in clinical use. They are inhibitors of squalene epoxidase an enzyme that is part of early stage of fungal cell membrane synthesis pathway. It prevents conversion of squalene to lanosterol, in turn ergosterol cannot be synthesized.

Fig 2: General mechanism of action of the antifungal agents. 12
6. Echinocandin: The class of echinocandin compounds is the fungal secondary metabolites, comprising a cyclic hexapeptide core with a lipid sidechain responsible for antifungal activity. The three commonly used echinocandin, anidulafungin, caspofungin and micafungin, inhibit the synthesis of cell wall β-1,3 glucan polysaccharides in the cell wall.

7. Sordarins: The sordarin family of compounds is characterized by a unique tetracyclic diterpene core. Sordarins selectively inhibit fungal protein synthesis by stabilizing the ribosome/EF2 complex. This unusual bioactivity makes sordarin a promising candidate for the development of new fungicidal agents.

1.6 Antifungal resistance: The development of drug resistance in fungi is a broad concept, which describes failure of current antifungal therapy to overcome the fungal infection. The antifungal therapies are designed to eradicate fungal infection by various mechanisms of action, like by disrupting their reproductive capabilities, destroying the cell walls, or by modifying the fungal DNA and altering the cell functioning. Antifungal resistance has been traditionally classified as [13-15].
a. Primary (intrinsic), present before the exposure to antifungals.
b. Secondary (acquired), developing after the exposure to antifungals owing to stable or transient genotypic alterations.
c. Clinical resistance, it can be classified as third type of antifungal resistance, which cover the progress or decline of antifungal properties. The macrolides were developed in 1952 having bacteriostatic properties. In 1956 Gold et al. reported the antifungal properties of the polyene amphotericin B, which was the first significantly effective systemic antifungal. It became the standard and soon it replaced hydroxy stilbamidine. Amphotericin B enjoyed the prime status of only antifungal agent available to treat systemic mycoses for nearly decade against which newer therapies for systemic mycoses were compared. In 1957 Fluocytosine, was developed as an antifungal agent who had failed to provide favorable results for the use as cytostatic agent. The use of fluocytosine as a monodrug often developed fungal resistance which led to use in combination with amphotericin B.

**Table 1**: Antifungal agents: Principal mode of action and resistance mechanism of fungal pathogens [15].

<table>
<thead>
<tr>
<th>Antifungal</th>
<th>Spectrum/comments</th>
<th>Mode of action</th>
<th>Mechanism of resistance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Polynes</td>
<td>Broad activity against Candida spp, Cryptococcus neoformans and filamentous fungi</td>
<td>Binding to Ergosterol</td>
<td>Alteration in ergosterol Biosynthesis</td>
</tr>
<tr>
<td>Amphotericin B</td>
<td></td>
<td>Impairment of nucleic acid biosynthesis</td>
<td>Decreased uptake of 5-FC.</td>
</tr>
<tr>
<td>Pyrimidine 5-fluorocytosine (5-FC)</td>
<td>Active against Candida spp and Cryptococcus spp.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Azoles</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1. Fluconazole</td>
<td>Active against Candida spp and Cryptococcus spp.</td>
<td>Inhibition of cytochrome P450 14α-lanosterol demethylase.</td>
<td>1. Upregulation of multi drug transporter genes. 2. Target alterations. 3. Alteration in Ergosterol biosynthetic pathway.</td>
</tr>
<tr>
<td>2. Itraconazole</td>
<td>Active against filamentous fungi.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4. Posaconazole</td>
<td>Related to itraconazole, but comparatively more active.</td>
<td>Inhibition of Squalene epoxidase</td>
<td>Unknown</td>
</tr>
<tr>
<td>Allylamines</td>
<td>Terbinafine Active against most dermatophytes.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Morpholines</td>
<td>Amorolfine Active against most dermatophytes</td>
<td>Inhibition of Sterol reductase and isomerase.</td>
<td>Unknown</td>
</tr>
<tr>
<td>Echinocandins</td>
<td>Caspofungin Active against Candida spp.</td>
<td>Inhibition of the cell wall synthesis</td>
<td>Unknown</td>
</tr>
</tbody>
</table>

Longer times for treatment with high relapse rate. This was followed by the commercial use of penicillin in the 1940s. In rapid succession, came the discovery and development of streptomycin in 1944 and benzimidazole the first azole to have notable antifungal activity was discovered in 1944. It was followed by the discoveries of chloramphenicol in 1947 and chlorotetracycline in 1948. In 1948 hydroxy stilbamidine, an antiprotozoal agent having antifungal action was used to treat blasticmycosis.

In 1951 Hazen and Brown, discovered the first polyene antibiotic called nystatin, commonly used topical and oral polyene. In 1952 substituted benzimidazole compounds were found to have antifungal properties. The macrolides were developed in 1952 having bacteriostatic properties.
to overcome the resistance. The first significant oral antifungal agent griseofulvin was developed in 1958, which became available for the treatment of superficial mycoses. Before griseofulvin, the treatment of superficial dermatophytoes was only by the topical drugs which were not especially effective against tinea capitis and onychomycosis. The semi-synthetic penicillin’s, cephalosporin’s and glycopeptides were developed from 1958 onwards. The development of chlorimidazole as a 5% cream in 1958, was beneficial over wide range of cutaneous mycoses. In the 1960s thiabendazole and mebendazole were reported to have antifungal and antihelminthic properties. In 1969 the imidazoles, clotrimazole and miconazole were introduced, which was soon followed by econazole in 1974. The allylamines discovered in 1974 are the other classes of antifungal agents that have a significant impact on antifungal therapy especially for superficial dermatomyces, including onychomycosis. Ketoconazole was developed in 1977 and since has become the standard among the azoles. In the mid-1980s two broad-spectrum, orally available triazoles, fluconazole (1982) and itraconazole (1984) were discovered. Between 1990-1999 to develop new antifungal agents and resulted into introduction of three azoles Voriconazole (2000)-Pfizer, Posaconazole (2005)-Schering-Plough, Ravuconazole (2007)-Bristol-Myers Squibb) and three echinocandins (Caspofungin (2002)-MSD, Anidulafungin(2004)-Astellas-Pfizer, Micafungin (2006)-Fujisawa) for their clinical use.

**Antifungal drug market: current & future scope**

11, 16, 17

**Current market scenario:** The antifungal drugs in the market are compromised of four established classes-polyenes (e.g. amphotericin B), azoles (e.g. fluconazole and itraconazole), allylamines (e.g. terbinafine) and the introduced echinocandins (e.g. caspofungin). Of these classes, only the first three are in use to treat systemic fungal infections. This market is predominated by the azoles, the most successful of which is Fluconazole, which was first launched in the mid-1980s by Pfizer pharmaceuticals, and is marketed as Diffucan. The toxicity problems of amphotericin B have been overcome to a little extent by the development of liposome based formulations, the most successful of which is Gilead’s Ambisome, but these formulations are generally very costly. Amphotericin was successfully replaced by fluconazole to treat invasive candida infections. Johnson & Johnson’s marketed product Sporanox (Itraconazole) is suffering in sales due to lack of new intravenous formulation. Pfizer’s Vfend (Voriconazole) and Schering’s posaconazole, the azoles with improved spectrum and activity will make further inroads into the amphotericin market. A new class of antifungal drug that has emerged recently is the echinocandins. The first representative of this is caspofungin, which is marketed by Merck as Candidas in 2002 for treatment of invasive Aspergillosis, having superior activity over amphotericin B. The second echinocandin; micafungin was launched as Funguard in 2002 by Fujisawa.

**Future market scenario:** Fungi are highly resistant eukaryotic microorganisms, and are becoming drug-resistant at an alarming rate to the approved compounds and some therapies later in clinical development. Despite the commercial success of the azoles, the drug interaction issues are a major impediment to the use of the azoles voriconazole, itraconazole and posaconazole. The interactions with cancer chemotherapy agents and immune suppressants are particularly difficult to handle clinically. On the other hand each year new pathogenic, mainly filamentous fungi are identified highlighting the need for broad spectrum coverage. Thus the rapidly increasing market of antifungal drugs have made the research and development of the newer and more effective antifungal agents of prime interest to both the industries and also to the academician around the globe.

**References**