



## Available classes

- Polyenes (cell membrane synthesis
- Azoles (cell membrane synthesis)
- Echinocandins (cell wall synthesis)
- Miscellaneous (nucleic acid, cell membrane synthesis)

#### Polyene mechanism of action

- Macrolide ring inserts into membrane parallel to phospholipid chains, binding to sterols
- Cylindrical channels form
- Cations, then macromolecules leak out
- Cell dies

# 1. Polyenes

- First antifungal antibiotics
- Isolated from <u>Streptomyces</u> spp.
- General structure:
  - Polyenes (multiple conjugated double bonds)
  - Macrolides (large rings with lactone linkage)



#### Polyene resistance

- Most clinically important fungi sensitive
- Dermatophytes resistant
- Inducible resistance rare (old drugs still work)
- Inherent resistance due to deminished membrane ergosterol with less affinity for drug

# Amphotericin B

- Colloidal dispersion in deoxycholate (bile salt)
- Protein bound. Urine and CSF concentrations low. Tissue stores slowly released
- Significant toxicity: – Infusion-related
  - Infusion-relate – Cumulative

## Polyenes: Nystatin

- 1950 in NYState
- Topical administration only
- Too toxic for systemic administration
- Uses:
  - Skin and mucosal candida infection especially oral thrush. No effect on dermatophtyes

# Infusion-related AmB toxicity

- Dramatic infusion-related fever, chills, nausea, vomiting, diarrhea, dyspnea
- ?cytokine/prostaglandin related
- Treatment: symptomatic premedication - Acetaminophen
  - Benadryl
  - Cortisone
  - Demerol
  - ?Duration of infusion

# Amphotericin B

- 1954 from Venezuela
- Not soluble in water at physiologic pH
- Not orally absorbed
- Occasional oral use of suspension for "topical" treatment of oral or esophageal candidiasis
- IV use: gold standard of antifungals

# Cumulative AmB toxicity

- **Renal**: characteristic cation-wasting nephropathy days-weeks into treatment. Low K+, Mg++, elevated creatinine. Treatment-limiting. (vasoconstriction, tubular cell lysis)
- Hematologic: characteristic normocytic anemia (direct marrow toxicity /renal)

# Amphotericin B uses

- Systemic fungal diseases caused by
  - Yeasts (candidiasis, cryptococcosis)
  - Molds (aspergillosis, mucormycosis)
  - **Dimorphs** (histo, blasto, cocci)
- Toxicity has shaped usage patterns





# Azole mechanism of action (and toxicity)

- Inhibit fungal cytochrome P450 enzymes which demethylate lanosterol to ergosterol
  - Block formation of ergosterol
  - Cause accumulation of toxic alpha-14 methyl esters in fungal cell
  - Sabotage membrane integrity
- "Fungistatic"

#### 2. Azoles

- 1970s to present
- From topical to powerful oral and IV drugs
- Imidazoles: 2N in 5-membered ring
- Triazoles: 3N in 5-membered ring

# Toxicity of Azoles

- inhibit cholesterol-dependent steroid hormone synthesis (testosterone; cortisol)
- Lead to ccumulation of metabolites with aldosterone-like effects
- Interfere with metabolism of other cytochrome P450 metabolized drugs

#### Resistance to Azoles

- Intrinsic, esp. nonalbicans Candida
- Inducible rare, but increasing with increasing use
  - Alteration in P450 enzymes
  - Membrane lipid changes with decreased permeability

# Newer azoles: Fluconazole

- 1990
- Soluble in water at neutral pH.
- Good oral absorption, urine and CSF penetration
- IV form available
- Toxicity primarily hepatic

#### Clincal uses:

- Cryptococcal meningitis
- Mucosal and esophageal candidiasis
- Systemic candidiasis (efficacy rivals AmB in some settings)
- Cocci

## Older Azoles

- Clotrimazole (Mycelex, Desenex, Lotrimin, Gynelotrimin)
- Miconazole (Monistat)
- Terconazole (Terazol)
- Topical onlyMinimal toxicity
- Minimal tox
  Used for
- dermatophyte and mucosal candidal infections

### Newer azoles: Itraconazole

- 1992
- Poorly watersoluble
- Protein and tissue-bound.
- Very high adipose and keratinized tissue levels

#### Clinical uses:

- Sporotrichosis
- Histoplasmosis
- Blastomycosis
- Cocci
  - Nail dermatophytes
  - Some activity against aspergillosis, sometimes.

# Newer azoles: Ketoconazole

- 1983
- Soluble in water at acid pH
- Highly protein/tissue bound
- Dose-related adrenal and testosterone suppression
- Clinical uses:
- Mucosal candidiasis (largely supplanted)
- Sporotrichosis
- Cocci
- Pityriasis and dermatophytes (Nizoral shampoo)

# Newest azole: Voriconazole

#### Clinical uses:

- Synthetic derivative of fluconazole with oral and IV dosing
   Enhance activity Asperg resistant
- Unique visual toxicity
- Enhanced in vitro activity against Aspergillus, resistant Candida
- Promising in vivo results

# 3. Echinocandins

- Inhibit fungal cell wall synthesis
- Irreversible inhibitors of 1,3 beta glucan synthase
- "Fungicidal" against wide range
- Little direct human toxicity

# Other agents

- **Griseofulvin** (1939) • Disrupts
- microtubulesActive only against
- dermatophytes, and not very.
- Relatively nontoxic
- Heading out
- thiocarbamatesInhibit squalene epoxidase

Allylamines and

- (ergosterol synthesis)
- Dermatophytes only
- Lamisil (terbinafine)



#### Fluorocytosine (5-FC, Flucytosine)

- Deaminated to 5-FU by bacterial and fungal cells
- Inhibits DNA synthesis in range of pathogens
- Rapid evolution of resistance precludes solo use
- Synergy in cryptococcisis , ?others
- Toxicity: **bone marrow suppression**, gastritis













