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## **Urinary Tract System**



## **Urinary tract infections(UTI's)**

- It is the 2<sup>nd</sup> most common infection (after RTI's).
- It is often associated with some obstruction of the flow of urine.
- It is more common in women more than men 30:1 (Why?).
- Incidence of UTI increases in old age(10% of men & 20% of women).

## What are the causes of UTI's

- Normally urine is sterile. Bacteria comes from digestive tract to opening of the urethra.
- Obstruction of the flow of urine(e.g. kidney stone)
- Enlargement of prostate gland in men(common cause)
- Catheters placed in urethra and bladder.
- Not drinking enough fluids.
- •Waiting too long to urinate.
- Large uterus in pregnant women.
- Poor toilet habits(wiping back to front for women)
- Disorders that suppress the immune system(diabetes & cancer chemotherapy).

**Bacteria responsible of urinary tract infections** 

- **Gm- bacteria (most common):**
- •E.coli (approx. 80% of cases)
- Proteus mirabilis
- •Klebsiella
- Pseudomonas aeruginosa
- **Gm+ bacteria :**
- •Staphylococcus Saprophyticus(Approx. 20%)

•Mycoplasma, Chlamydia trachomatis & N. gonorrhea (limited to urethra, unlike E.coli may be sexually transmitted)

### **Urinary tract infections can be:**

### •Simple:

Infections do not spread to other parts of the body and go away readily with treatment ( Due to E.coli in most cases).

#### •Complicated:

Infections Spread to other parts of the body and resistant to many antibiotics and more difficult to cure.{Due to hospitalacquired bacteria(E.coli, Klebsiella,, Proteus, Pseudomonas, enterococci, staphylococci)}

## **Treatment of UTI's**

#### **Antibiotics:**

- Co-trimoxazole(SMX/TMP) ),p.o.
- Nitrofurantoin,p.o.
- Tetracyclines, e.g. Doxycycline, p.o.
- Aminoglycosides, e.g. gentamicin
- cephalosporins(e.g.ceftriaxone&ceftazidime
- Quinolones, e.g. ciprofloxacin,p.o.

## Co-trimoxazole (Bactrim, Septra) Sulfamethoxazole- Trimethoprim (SMX) (TMP)

Alone, each agent is bacteriostatic

- Together they are bactericidals(synergism)
- The optimal ratio of TMP to SMX in vivo is 1:20

(formulated 5(SMX):1(TMP); 800mg SMX+160mg TMP; 400 mg SMX+ 80 mg TMP; 40 mg SMX+8 mg TMP).

### **MECHANISM OF ACTION**



#### Absorption, metabolism& Excetion

#### **Sulfonamides**

Mainly given orally

Rapidly absorbed from stomach and small intestine.

Widely distributed to tissues and body fluids (including CNS, CSF), placenta and fetus.

Absorbed sulfonamides bind to serum protein(approx. 70%). Metabolized in the liver by the process of acetylation. Eliminated in the urine, partly as such and partly as acetylated derivative.

#### Trimethoprim (TMP)

Usually given orally, alone or in combination with SMX Well absorbed from the gut Widely distributed in body fluids & tissues (including CSF) More lipid soluble than SMX Protein bound (approx.40 %) 60% of TMP or its metabolite is excreted in the urine **TMP concentrates in the prostatic fluid**.

#### **ADVERSE EFFECTS**

- 1.Gastrointestinal-Nausea, vomiting
- 2. Allergy
- 3. Hematologic
  - a) Acute hemolytic anemia
    - a) hypersensitvity b) G6PD deficiency
  - b) Megaloblastic anemia due to TMP.
- 4. Drug interactions
  - Displace bilirubin- if severe kernicterus Potentiate warfarin, oral hypoglycemics.

### **CONTRAINDICATIONS**

- 1. Pregnancy
- 2. Nursing mother
- 3. Infants under 6 weeks
- 4. Renal or hepatic failure
- **5. Blood disorders**

### Nitrofurantoin

#### **Antibacterial Spectrum:**

Effective against E. coli and Staph. saprophyticus, but other common UT gm- bacteria may be resistant.

#### **Mechanism of action of nitrofurantoin**

Sensitive bacteria reduce the drug to an active agent that inhibits various enzymes and damages DNA.

#### **Pharmacokinetics of nitrofurantoin**

- •Absorption is complete after oral use
- Metabolized (75%)& excreted so rapidly that no systemic antibacterial action is achieved.
- Concentrated in the urine(25% of the dose excreted unchanged)
- •It turns urine to a dark orange-brown.

### **Adverse effects of nitrofurantoin**

GI disturbances: bleeding of the stomach, nausea, vomiting and diarrhea(must be taken with food). Headache and nystagmus. Hemolytic anaemia(G6PD deficiency)

#### **Containdications:**

Pts with G6PD deficiency

Neonates

Pregnant women(after 38 wks of pregnancy)

#### **Therapeutic Uses of nitrofurantoin**

- It is used as urinary antiseptics . Its usefulness is limited to lower UTI's & cannot be used for upper UT or systemic infections.
- Dose: 50-100 mg, po q 6h/7 days.
- Long acting: 100mg twice daily.

## Tetracyclines (e.g. Doxycycline)

It is a long acting tetracycline Mechanism of action Inhibit protein synthesis by binding reversibly to 30 s subunit

# **Doxycycline (Cont.)**

#### **Pharmacokinetics**

- Usually given orally
- Absorption is 90-100%
- Absorbed in the upper s. intestine & best in absence of food
- Food & di & tri-valent cations (Ca, Mg, Fe, AL) impair absorption
- **Protein binding 40-80 %**
- **Distributed well, including CSF**
- **Cross placenta and excreted in milk**
- Largely metabolized in the liver

# **Doxycycline (Cont.)**

#### Side effects

- 1. nausea, vomiting ,diarrhea & epigastric pain(give with food)
- 2. Thrombophlebitis i.v
- 3. Hepatic toxicity ( prolonged therapy with high dose )
- 4. Brown discolouration of teeth children
- 5. Deformity or growth inhibition of bones children
- 6. Phototoxicity
- 7. Vertigo
- 8. Superinfections.

## **Contraindications of doxycycline**

• Pregnancy

• Breast feeding

Children(below 10 yrs)

### **Therapeutic Uses of Doxycycline**

- •Treatment of UTI's due to Mycoplasma & Chlamydia, 100 mg p.o bid for 7 days.
- Prostatitis

## Aminoglycosides

#### e.g. GENTAMICIN, i.m, i.v.

- Bactericidal antibiotics
- Inhibits protein synthesis by binding to 30S ribosomal subunits.
- Poorly absorbed orally(highly charged).
- Only active against gram negative aerobic organisms.
- cross placenta.

## **Gentamicin(CONT)**

- Excreted unchanged in urine
- More active in alkaline medium
- Adverse effects :
- Ototoxicity
- Nephrotoxicity
- Neuromuscular blocking effect

#### **Therapeutic uses of Gentamicin in UTI's**

• Severe infections caused by gram negative organisms (pseudomonas or enterobacter).

**Cephalosporins** (Detail was explained in respiratory lec.) **3**<sup>rd</sup> generation cephalosporins **Ceftriaxone & Ceftazidime** Mainly effective against gm-bacteria. Acts by inhibition of cell wall synthesis **Bactericidal** They are given parenterally

Given in severe / complicated UTIs

& acute prostatitis

## Fluroquinolones

(Detail was explained in respiratory lec.)

#### e.g. ciprofloxacin

Active against gram negative aerobic organisms. Mechanism of action

• Inhibits DNA gyrase enzyme

#### **Clinical use**

- UTIs caused by multidrug resistance organisms as pseudomonas.
- Prostatitis ( acute / chronic )

### **Adverse effects**

- Nausea , vomiting , diarrhea
- CNS effects ( confusion, insomnia, headache, anxiety).
- Damage of growing cartilage(arthropathy)
- Phototoxicity(avoid excessive sunlight)