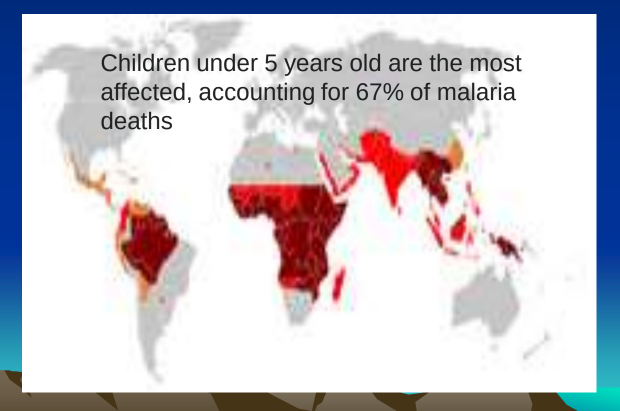
# **Antimalarial Drugs**

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#### **Malaria**

 In 2019, there were an estimated 229 million cases of malaria(296 million in 2015) which resulted in an estimated 409,000 deaths.



### **Antimalarial Treatment**

• SuppressiveTreatment (المعالجة القمعية) = Clinical Cure: Chloroquin, Quinine, Quinidine, Doxycyline, Clindamycin, Mefloquine, and Halofantrine.

• Radical Cure(المعالجة الجذرية): Chloroquin followed by Primaquine, required for *P vivax and P ovale*.

Prophylaxis: Chloroquin, Mefloquin, "Malarone", and Doxycycline.

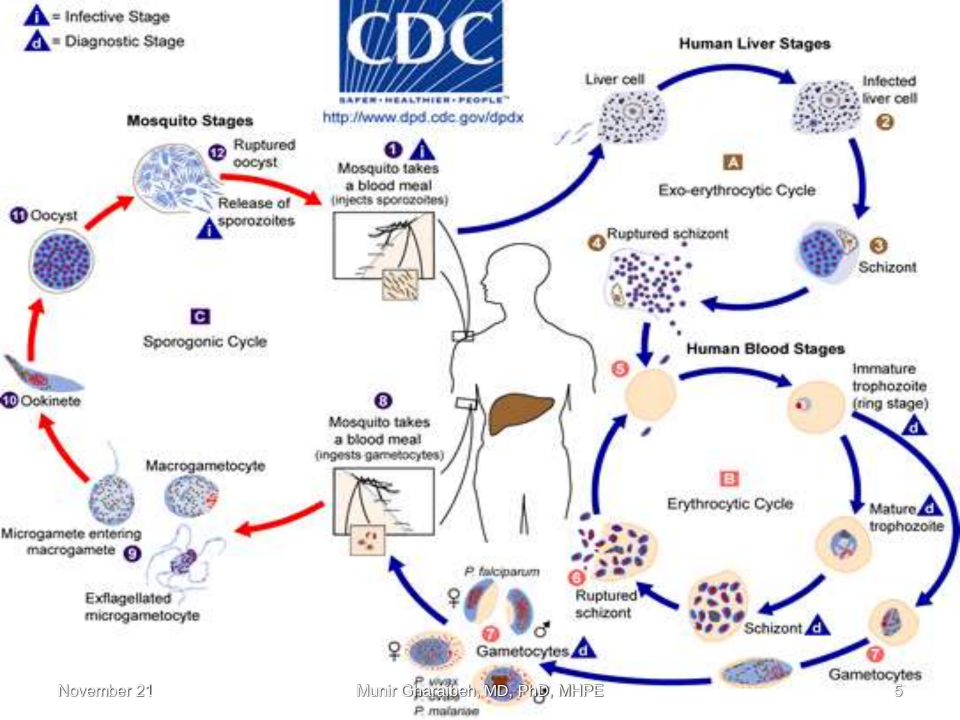
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#### **Malarial Parasites**

Plasmodium falciparum

(only erythrocytic, serious, resistance).

- Plasmodium vivax.
- Plasmodium malariae.
- Plasmodium ovale.



#### Synthetic 4-Aminoquinolone

Specific uptake mechanism is present in the parasite, the drug accumulates in the parasite to inhibit polymerization of heme into hemozoin and thus parasite is poisoned by heme.

Well absorbed, distributed, bound to tissues.

- Schizonticide for all four types of malaria.
- Drug of choice in the treatment of nonfalciparum and sensitive falciparum malaria.
- Does not eliminate dormant liver forms of *P. vivax* and *P.* ovale, so, Primaquine must be added for their radical cure.

#### Resistance:

Very common with *P. falciparum* and increasing with *P.vivax*.

Due to mutation in P170 glycoprotein (PfCRT) works as a drug-transporting pump mechanism.

- Very practical, convenient(oral), rapid action, low cost, and safe.
- Started immediately after diagnosis.
- Other doses are given after 6 hours, 24 hours and last dose after 48 hours.
- However, does not eliminate dormant liver forms of P.vivax and P.ovale.

#### Also effective in:

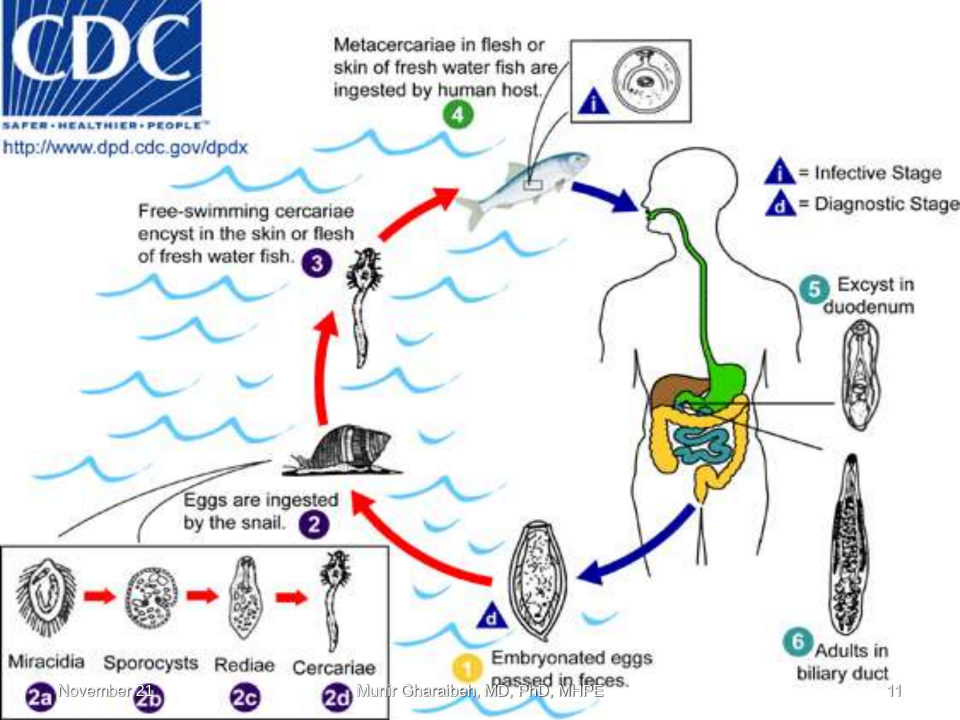
Rheumatoid arthritis.

LE.

Amebic liver abscess.

Photoallergic reactions.

Clonorchis sinensis.



#### **Side Effects:**

Headache, dizziness,
Itching and rash,
Nausea, vomiting, anorexia
Unmasking of LE, psoriasis and porphyria.
Corneal deposits, blindness, blurring of vision,

### Quinine(1820) and Quinidine

Cinchona tree from South America.



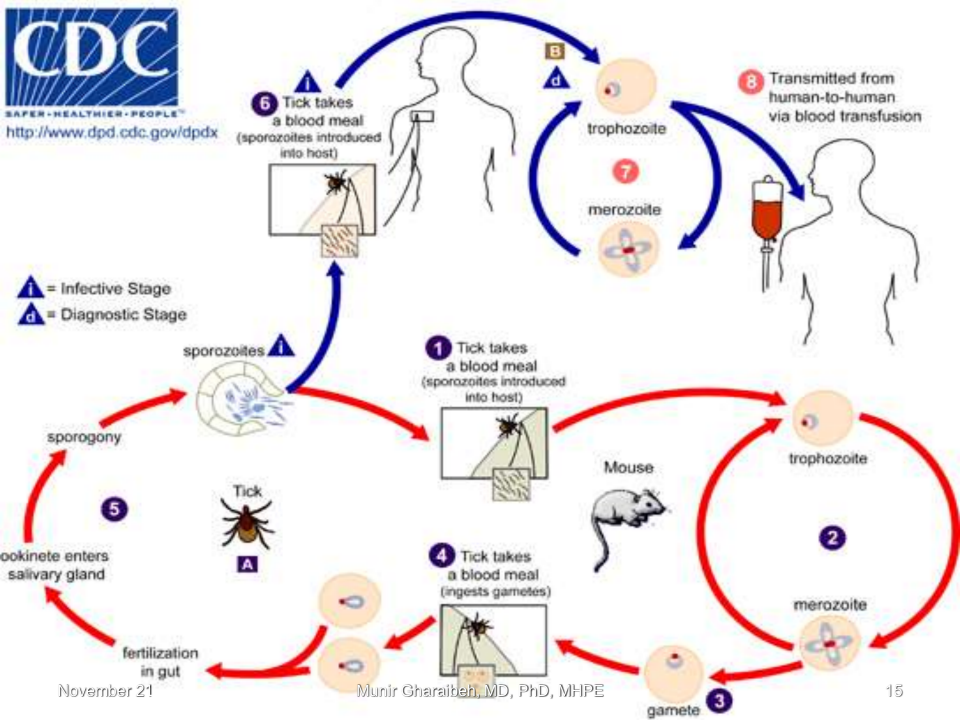
General protoplasmic poison: will affect the feeding mechanism of the parasite.

Resistance is uncommon.

Effective rapid schizonticide therapy for severe falciparum, chloroquine-resistant malaria, usually in combination with another drug (e.g. Doxycycline or Clindamycin) to shorten duration of use.

### Quinine and Quinidine

- Also effective for Babesia microti infection.
- Also, for nocturnal leg muscle cramps (Arthritis, DM, thrombophlebitis, arteriosclerosis, varicose veins)



### Quinine and Quinidine

#### **Adverse Effects:**

Cinchonism: Tinnitus, headache, nausea, dizziness, flushing, visual disturbances. Later, auditory abnormalities, vomiting, diarrhea, and abdominal pain.

**Blood dyscrasias.** 

Hypersensitivity, hypoglycemia, uterine contractions.

Hypotension, QT prolongation.

Blackwater fever (hemolysis, hemoglobinemia,

### Mefloquine

Blood schizonticide, not for liver forms.
 Used for resistant P. falciparum (single oral dose).

Also for suppressive and prophylactic treatment (weekly doses).

- Nausea, vomiting, diarrhea, pain.
- Vertigo, dizziness, headache, rashes and visual alterations.
- Psychosis, hallucinations, confusion, anxiety, depression.

# **Primaquin**

8-aminoquinolone
 Unknown mechanism.
 Drug of choice; the only available one, for eradication of exoerythrocytic forms of malaria after treatment with chloroquin.

Hemolysis in G6PD deficient patients. Also, nausea, distress, headache, pruritis, leukopenia and agranulocytosis.

#### **Atovaquone and Proguanil**

- Usually in fixed combination = "Malarone".
- Recommended drug for prophylaxis.
- Atovaquone also approved for *P. jiroveci* pneumonia, although has lower efficacy than
   Trimethoprim-sulfamethaxazole combination.
- Can cause fever, rash, nausea, vomiting, diarrhea, headache, and insomnia.

#### **Pyrimethamine**

- **Inhibits DHF Reductase**
- Slow and long acting drug.
- Effective on erythrocytic forms of all species.
- Not for severe malaria.
- Preferential binding to parasitic enzyme.
- Usually combined with Sulfadoxine" Fansidar" or Sulfones which inhibit Dihydropteroate synthase.
- No longer recommended for prophylaxis.
- Also, for Toxoplasmosis( in higher doses ),
- and P. jeroveci.

# **Pyrimethamine**

#### **Adverse Effects:**

Anorexia, Vomiting, Leucopenia, Thrombocytopenia, glossitis

**CNS: Stimulation, Convulsions** 

Allergic reactions including Stevens-Johnson

**Syndrome** 

#### **Antibiotics**

- Tetracycline.
- Doxycycline.
- Clindamycin.
- Azithromycin.
- Fluoroquinolones.

Active against erythrocytic forms of all species. Usually for chloroquine-resistant strains. Also effective against other protozoal diseases.

#### **Halofantrine and Lumefantrine**

Rapidly effective against erythrocytic forms of all species.

Usually for chloroquine-resistant strains.

Well tolerated, except for cardiac toxicity (QT prolongation)

# **Artemisnin= Qinghaosu**

- Artesunate.
- · Artemether.



- Derivatives of Artemisia(الشيح) used by Chinese since 2000 years.
- Rapidly acting schizonticides against all species.
- No documented resistance.
- Work by free radical formation or ATP inhibition.
- Only drugs reliably effective against quinineresistant and multi-drug resistant strains.
- High cost.
- N.V.D. and neurotoxicity in animals.