



Sofia Zago died of cerebral malaria after developing a fever & slipping into a coma last September in north Italy

#### Sofia had NOT travelled to any at-risk countries

# World Health Organization declared Italy free of malaria in 1970



# HOW DID SOFIA CONTRACT MALARIA?

©Could it be transmitted from **immigrants**? Malaria is not transmitted from person to person

Could **climate** change to blame for the first-home grown case in 50 years?

With global climate change, the potential for the reappearance of malaria in countries where it was previously eradicated exists, but is relatively small.

Could a **re-used needle** at the hospital where she was treated to blame?

Could the disease-carrying mosquitoes arrived by plane in people's suitcases?





Classify the main <u>antimalarial drugs</u> depending on their goal of therapy

Optimize the pharmacokinetics & dynamics of main drugs used to treat attack or prevent relapses

State the WHO therapeutic strategy for treatment

Int on the CDC recommendations for prophylaxis in travelers to endemic areas.



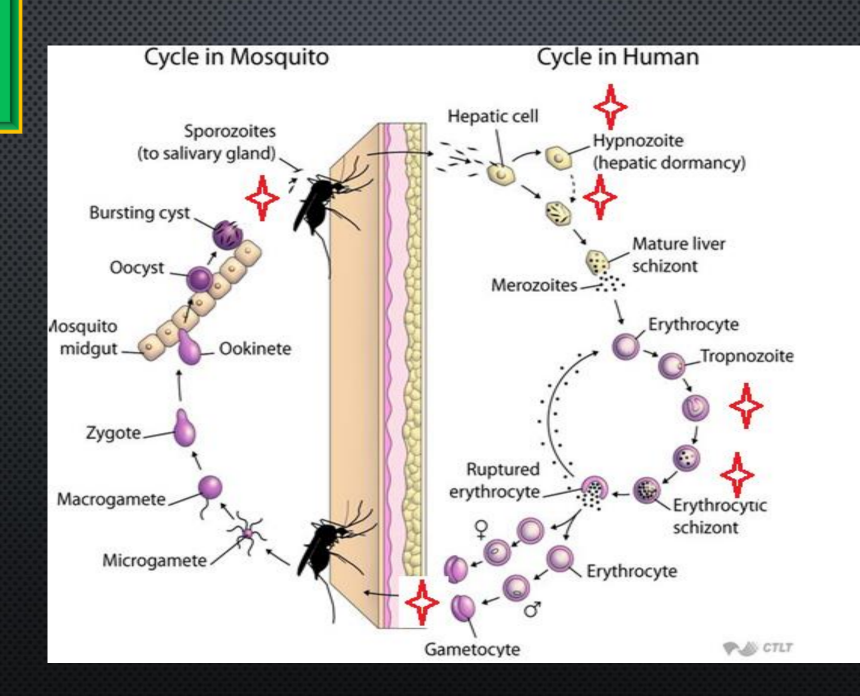
#### According to WHO:

212 million cases of malaria worldwide in 2015 & 429,000 deaths. 90% of malaria cases & deaths occur in **Africa**. Children under 5 are most at risk.

**Four** species of plasmodium typically cause human malaria:

- Plasmodium falciparum,
- P vivax,
- P malariae, and
- P ovale.

#### Cycle & Drugs site of action





# THERAPEUTIC CLASSIFICATION:

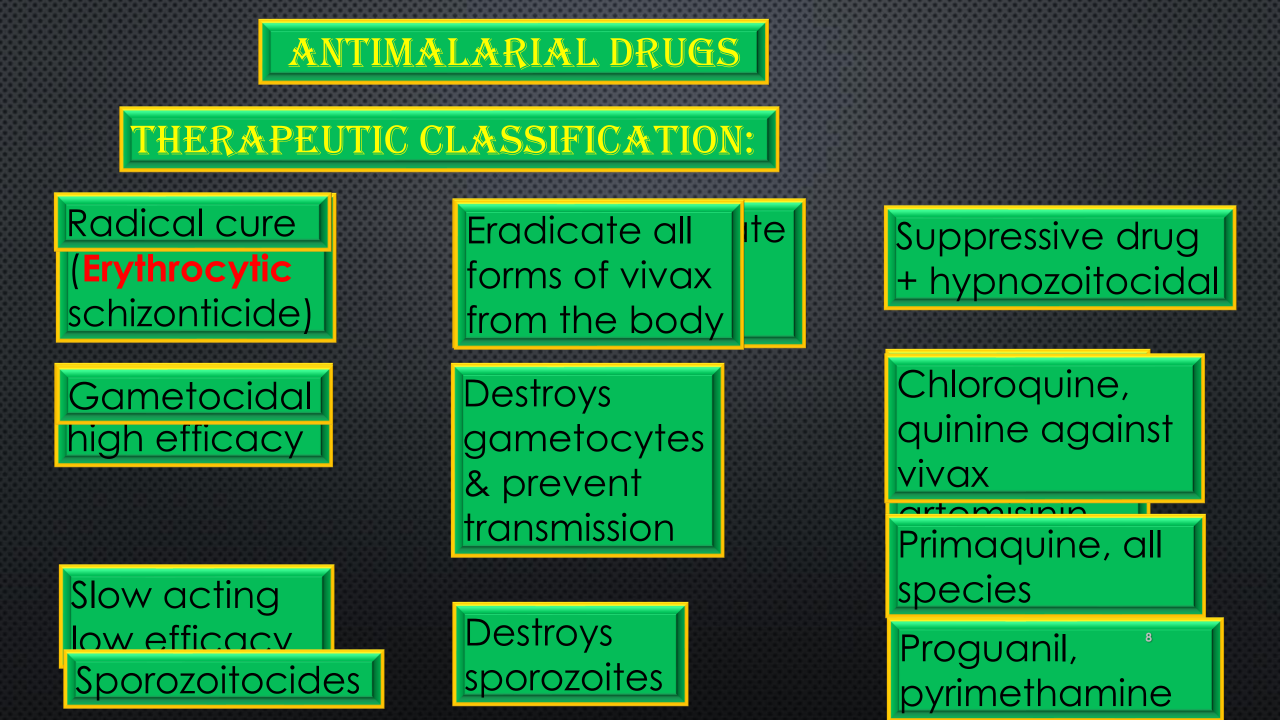
## Causal prophylaxis

Destroys parasite in **liver** cells & prevent invasion of erythrocytes





Suppresses the **erythrocytic** phase & thus attack of malaria fever Chloroquine, mefloquine, doxycycline





# Artemisinin is the active principle of the plant Artemisia annua (qinghaosu)

Fast acting blood Schizontocide

Affect all forms including multidrug resistant P. falciparum

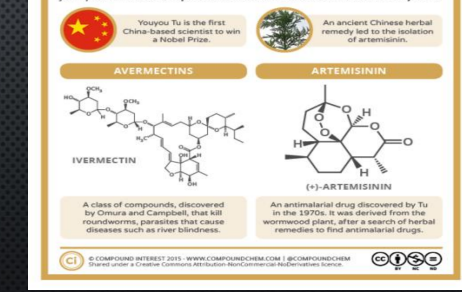
#### Short duration of action

e High recrudescence rate after short-course therapy



#### **NOBEL PRIZE IN MEDICINE 2015**

The Nobel Prize in Physiology or Medicine 2015 was awarded with one half jointly to William C. Campbell & Satoshi Omura & the other half to Youyou Tu.



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Poorly soluble in water & oil, can only be used orally.

#### PHARMAKOKINETICS

### Rapidly biotransformed in liver into di-hydroartesiminin + active metabolite

#### @Artemisinin, artesunate, artemether are prodrugs

Derivatives are rapidly absorbed orally & Widely distributed

Artemisinin  $t^{1/2} \rightarrow 4$  hrs

**@Artesunate** t½ 45 min (water-soluble; oral, IV, IM, rectal administration) **@**Artemether t½ 4-11 hrs, (lipid-soluble; oral, IM, & rectal administration). Induce its own CYP-mediated metabolism→  $\uparrow_{0}$ clearance 5 fold.

# ARTEMESININ & ITS DERIVATIVES

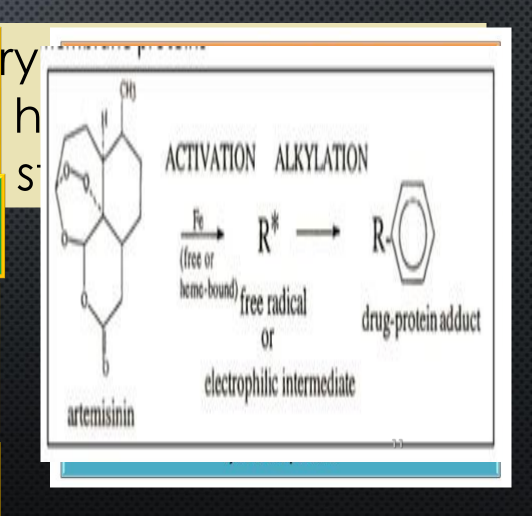
# MECHANISM

 They have endoperoxide bridges
 Haem iron cleaves this bridge to yield carboncentered free radicals in parasite, that will →

@Alkylate membranes of parasite's food vacuole & mitochondria > no energy

Irreversibly bind & inhibit sarco-endoplasmic reticulum Ca<sup>2+</sup>-ATPase of the parasite, thereby inhibiting its growth

Inhibiting formation of transport vesicles + no food vacuoles.





#### CLINICAL USES

Because artemisinin derivatives have short t<sup>1</sup>/<sub>2</sub>,
 (1) Monotherapy should be extended beyond disappearance of parasite to prevent recrudescence or

(2) by combining the drug with long- acting antimalarial drugs (Ex. mefloquine)

#### PREPARATIONS

Artesunate IV or IM preparations for severe complicated cases as cerebral malaria (24 h) followed by complete course of ACT.



# PREPARATIONS

#### @Artemisinin-based combination therapies (ACTs):

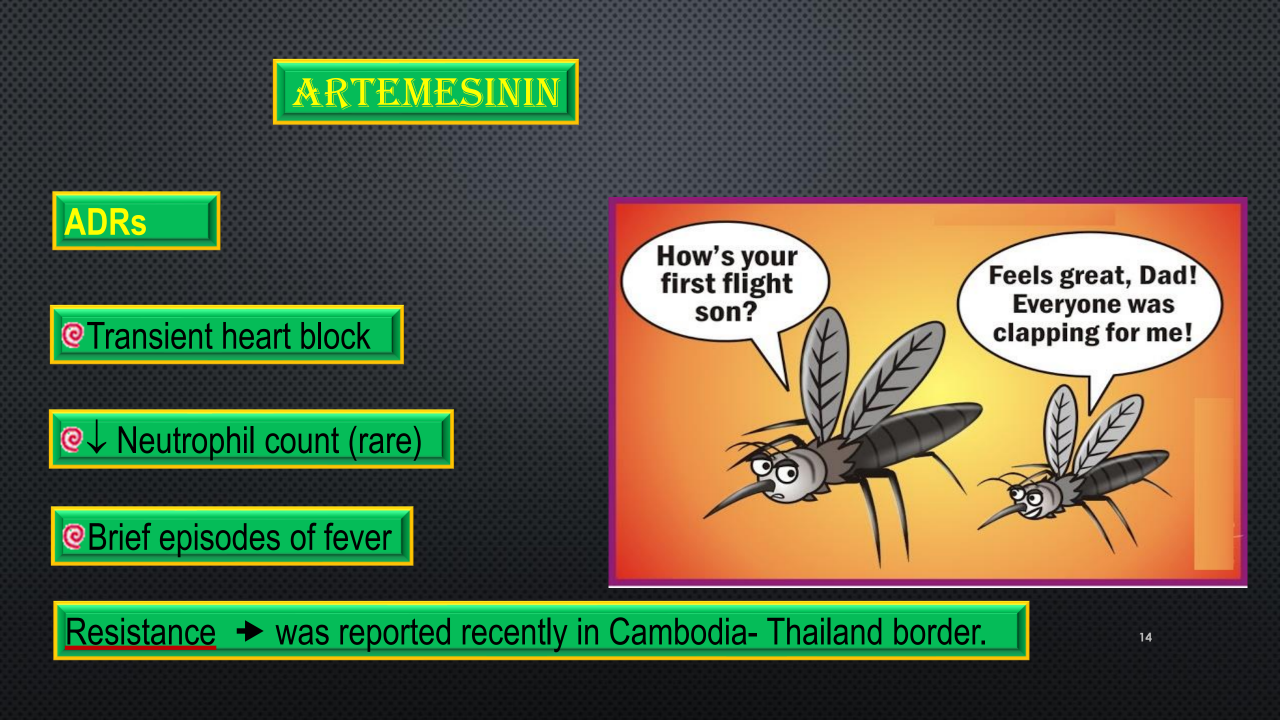
Artemether + lumefantrine

Artemether + amodiaquine

Artemether + mefloquine

>Artemether + sulfadoxine- pyrimethamine.

**CDC/Jahles Gathany** 





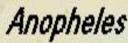


#### Potent **blood** Schizontocide

# Active against all forms of the schizonts (exception is chloroquine-resistant P.f. & P.v.)

Not active against tissue shizonts

Gametocide:-Against all species except P. falciparum.



(from Iowa State Univ. Entomology Image Gallery)

CLOBAL

250 mg

Chloroquine Phosphate Tablets, USP

## CHLOROQUINE

## PHARMACOKINETICS

Rapidly & completely absorbed from the GIT, given po
 Has high volume of distribution (100-1000 L/kg);
 Released slowly from tissues & metabolized in liver

Concentrated into parasitized RBCs

@ Excreted in the urine 70% unchanged @ Initial  $t\frac{1}{2}$  = 2-3 days & terminal elimination  $t\frac{1}{2}$ =1-2 months.

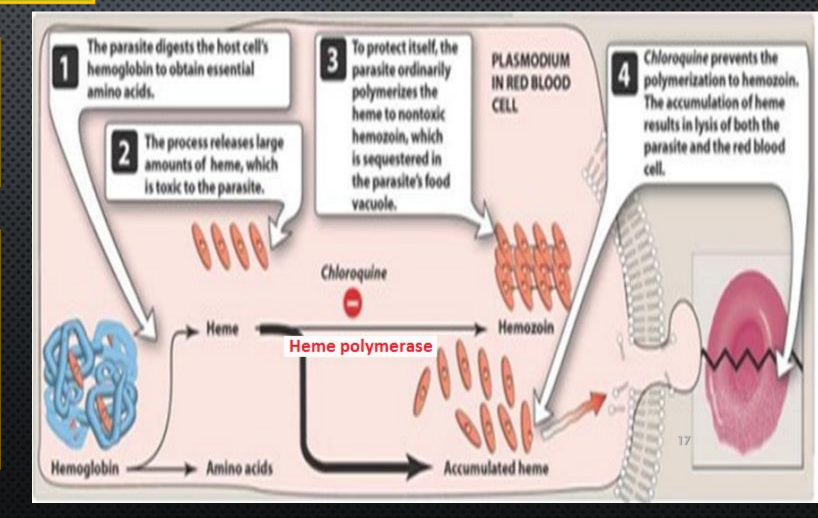


# CHLOROQUINE

## **MECHANISM OF ACTION**

<u>Malaria Parasite</u> digest host cell's Hb to utilize globin & obtain amino acids

Heme is released → Toxic
So parasite detoxifies it by
heme polymerase →
Hemozoin (NonToxic) & traps
it in food vacuoles.

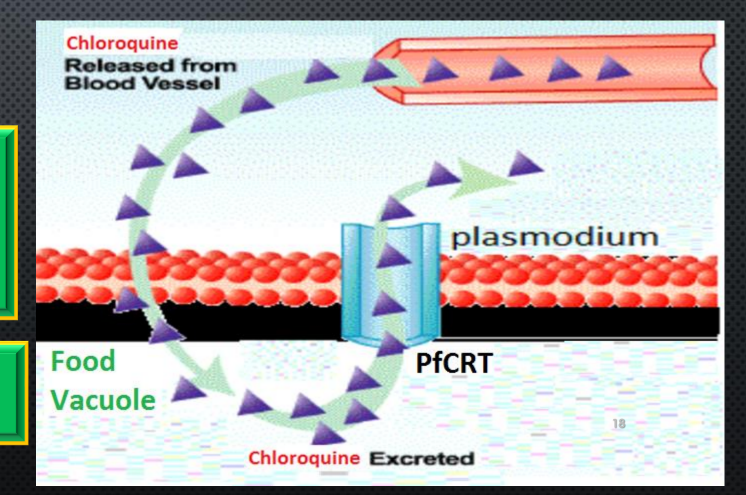




# RESISTANCE

Resistance against the drug develops as a result of <u>mutation</u> of the chloroquine resistance transporter (PfCRT)

PfCRT enhances the efflux of chloroquine from the food vacuole.



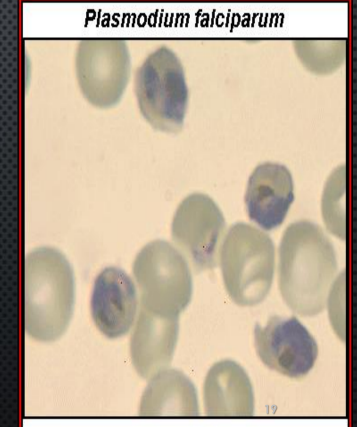


#### THERAPEUTIC USES

Used to eradicate **blood** schizonts of *Plasmodium*. It is given in loading dose to rapidly achieve effective plasma conc.



Rheumatoid arthritis.



(original image provided by Steve Aley)

# CHLOROQUINE



- 1. Mild headache & visual disturbances
- 2. GIT upsets; Nausea, vomiting
- 3. Pruritus, urticaria.

Prolonged therapy & high doses:

 Ocular toxicity: Loss of accommodation, lenticular opacity, retinopathy

 Ototoxicity

 Weight loss

 Bolus injection → hypotension & dysrrhythmias

 @Safe in pregnancy

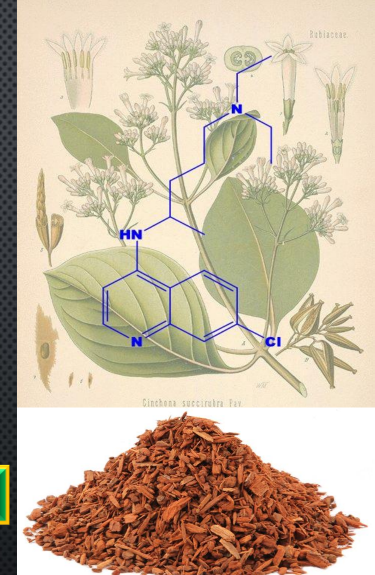


The main alkaloid in cinchona bark

Potent blood Schizontocide of ALL malarial parasites & gametoside for P vivax & ovale but not falciparum. It is Not active against liver stage parasites.

Output Depresses the myocardium, reduce excitability & conductivity

Mild analgesic, antipyretic, stimulation of uterine smooth muscle, curare mimetic effect.





# PHARMACOKINETICS

#### Rapidly & completely absorbed from the GIT

Peaks after 1-3 hours

Metabolized in the liver & excreted in urine

@ 5-20% excreted in the urine unchanged

0 t<sup>1</sup>/<sub>2</sub> = 10 hrs but longer in sever falciparum infection (18 hrs)

Administered: orally in a 7 day course or by slow IV for severe *P. falciparum* infection.



# MECHANISM Same as chloroquine

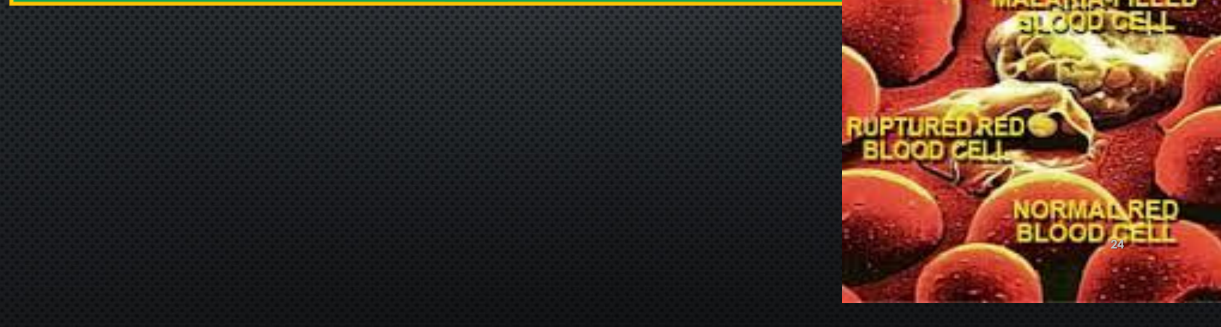
MECHANISM OF RESISTANCE Like chloroquine, by mutation of chloroquine resistance transporter, also increased expression of **P-glycoprotein transporter**.





## CLINICAL USES:

# Parenteral treatment of severe falciparum malaria Oral treatment of falciparum malaria Nocturnal leg cramps.





# ADRS

#### With therapeutic dose + poor compliance + bitter taste

Higher doses + @ Cinchonism + *(tinnitus, deafness, headaches, nausea & visual* 

@Abdominal pain & diarrhea
 @Rashes, fever, hypersensitivity reactions

Output 
Out

Safe in pregnancy

Blood dyscarasis; anaemia, thrombocytopenic purpura & hypoprothi

@Blackwater fever, a fatal condition in which acute haemolytic anae with renal failure due to a hypersensitivity reaction to th

IV → neurotoxicity → tremor of the lips & limbs, delirium, fits, stimulation followed by depression of respiration & coma.



# CONTRAINDICATIONS

#### Prolonged QT Interval

Glucose-6-Phosphate Dehydrogenase deficiency & pregnancy

Myasthenia Gravis

Output 
Out

Optic Neuritis, auditory problems

Ose should be reduced in renal insufficiency.

Glucose-6-phosphate G-6-P-D enzyme 6- Phosphoguconate + NADPH+ $H^+$ In case of G-6-P-D Leads to Inadequate supply of NADPH Leads to Reduced level of Glutathione Glutathione protects Hb oxidation



# DRUG INTERACTIONS

 Antacids: Antacids containing aluminum &/or magnesium may delay or decrease absorption of quinine

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• Mefloquine

# • Quinine can raise plasma levels of warfarin & digoxin.

## PRIMAQUINE

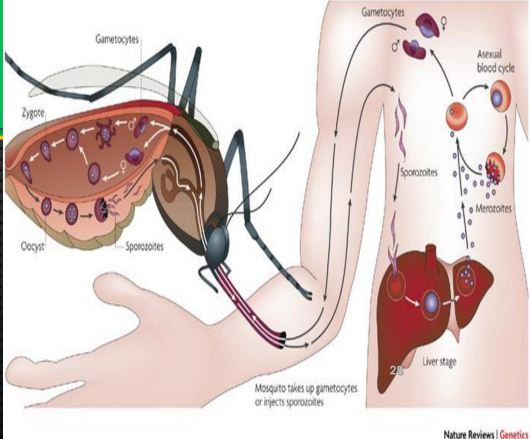
Pypnozoitocides against liver hypnozoites
gametocytocides against the 4 human malaria species
Radical cure of *P. ovale & P. vivax*Prevent spread of ALL forms (chemoprophylaxis)

#### PHARMACOKINETICS

Well absorbed orally

- Rapidly metabolized to etaquine & tafenoquine
- more active forms
- @ t½ → 3-6 h.









Not well understood, It may be acting by:-

 @Generating ROS → can damage lipids, proteins & nucleic acids in the parasite
 @Interfering with the electron transport → no energy

Inhibiting formation of transport vesicles + no food vacuoles

Resistance; 

Resistance; 

Rare when primaquine & chloroquine are combined.

Primaquine

Converted to electrophiles

Generates reactive oxygen species

- Interferes with oxygen transport system



ANTIMALARIAL DRUGS



# Radical cure of relapsing malaria, 15 mg/day for 14 days

In falciparum malaria: a single dose (45 mg) to kill gametes & cut down transmission

G-6-PD NORMAL	15mg per day x 14
G-6-PD deficiency (Mild African form)	45mg per week for 8
G-6-PD deficiency (More severe Mediterranean variety)	30mg per week for 30 weeks
o fotus is	the urine

Primaguine

Should be avoided in pregnancy (the fetus is relatively G6PD-deficient & thus at risk of hemolysis) & G6PD deficiency patients



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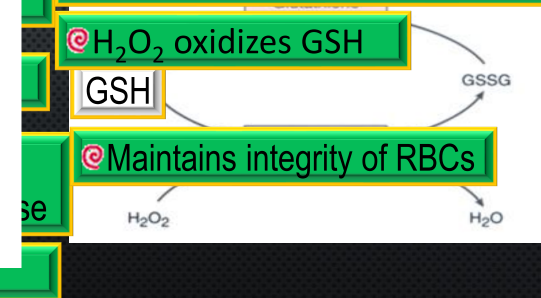
# ADRS

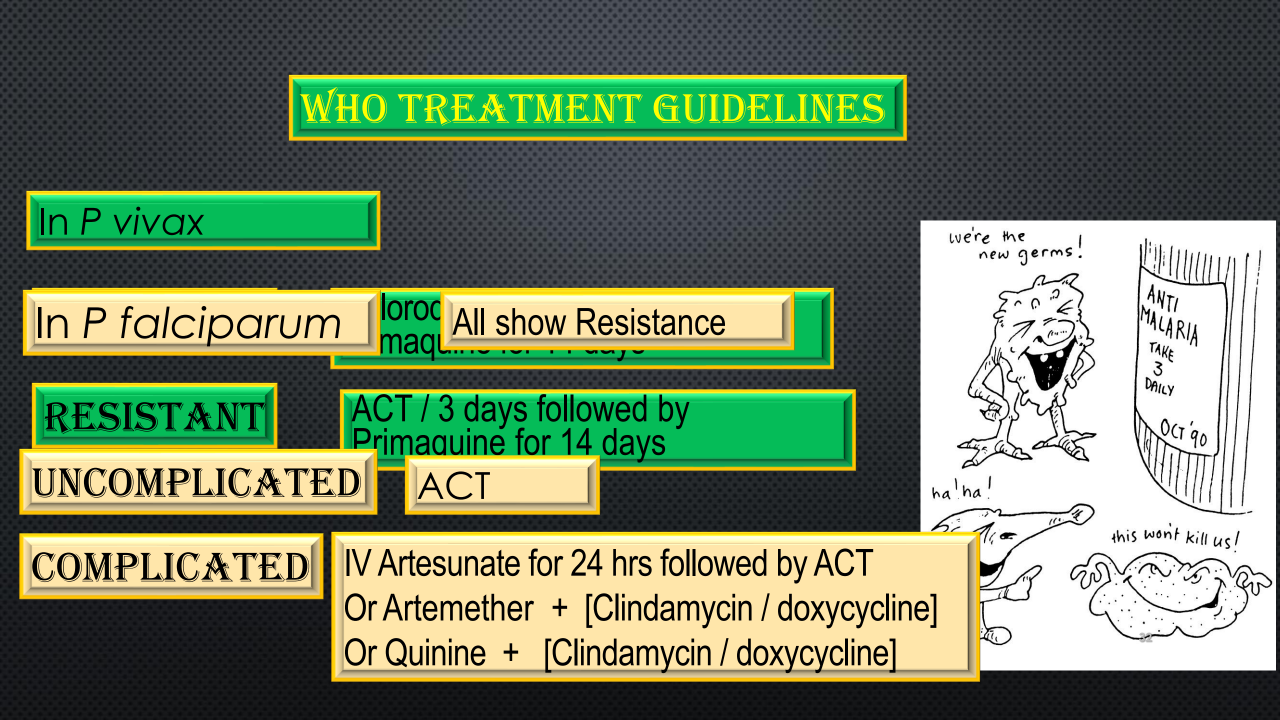
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# <u>At regular doses</u> → patients with G-6-PD deficiency → hemolytic anemia.

Oxidation of primaqune produces free radicals

@Free radicals will cause oxidative
damage of RBCs →Hemolysis







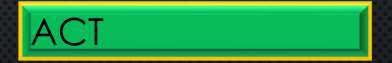
# IN FALICPARUM

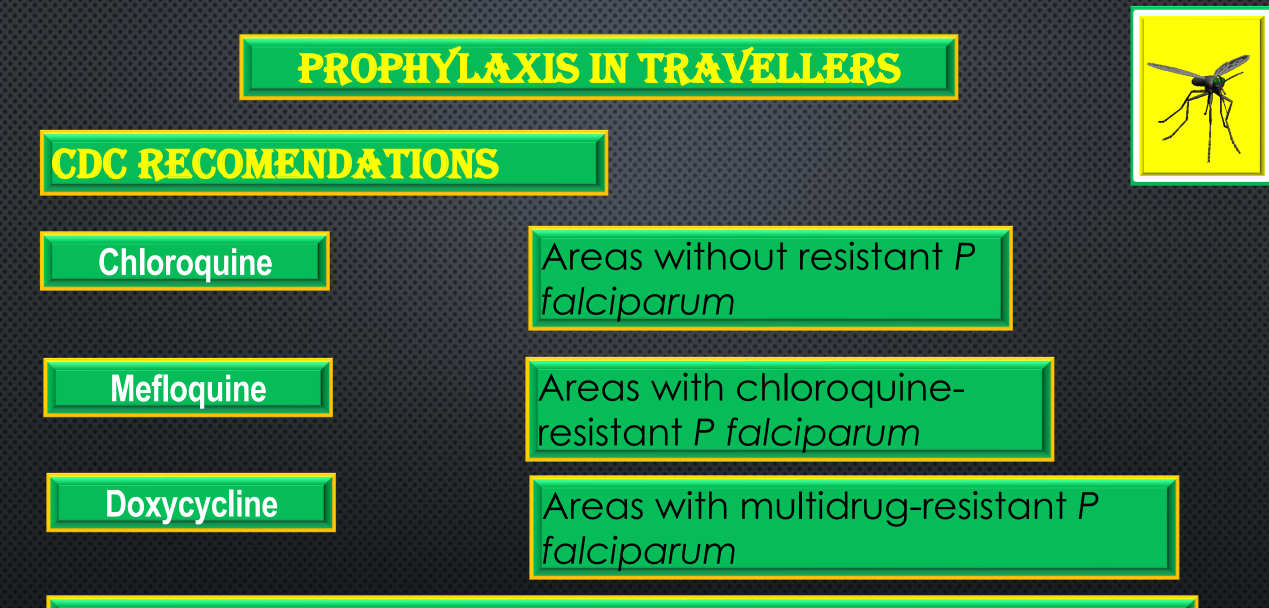
# SPECIAL RISK GROUPS

Pregnancy; 1<sup>st</sup> trimester

Pregnancy; 2<sup>nd</sup> & 3<sup>rd</sup> trimester Lactating women Infants & young children







Begin 1-2 weeks before departure (except for doxycycline 2 days) & continue for 4 weeks after leaving the endemic area.

# THANK U....