

Life Cycle

Has 2 hosts: vertebrate (**intermediate**) and mosquito (**definitive**)

Definitive Host: Intermediate
Sporulating Host: Asexual
Cycle (Sporogony) Cycle (Schizogony)

EXO-ERYTHROCYTIC CYCLE

1. Anopheles mosquito (definitive host), inoculates **sporozoites** into human host (intermediate)

Sporozoites are the infective stage to man

2. Sporozoites infect **liver parenchymal cells** and mature into **schizonts**, which produce merozoites

P. vivax and *P. ovale* assume a dormant stage (**hypnozoites**), and can persist in the liver

3. Infected cell ruptures and releases **merozoites**

ERYTHROCYTIC CYCLE

4. Merozoites infect **RBCs** and become **trophozoites** (ring stage)

Merozoites are the infective stage for RBCs

Life Cycle (cont)

5. Trophozoites mature into **schizonts**, which rupture and release **merozoites**

Some differentiate into **gametocytes**, which are ingested by the mosquito looking for a blood meal

6. Ruptured schizonts are responsible for the **clinical manifestations** of the disease

SPOROGENIC CYCLE

7. The **gametocytes** are ingested by an anopheles mosquito and make their way into the **stomach** of the mosquito

8. While in the stomach, the **microgametes** (male) penetrate the **macrogametes** (female), which generate **zygotes**

Microgametocyte exflagellates and produces 8 sperm-like microgametes

9. The zygotes become **motile** and **elongated**

These are called **ookinetes**

10. The ookinetes invade the **midgut wall** of the mosquito, where they develop into **oocysts**

Life Cycle (cont)

11. The **oocysts** grow, rupture, and release **sporozoites**, which makes their way into the **salivary glands** of the mosquito

12. Inoculation of the sporozoites into a new human host perpetuates the malaria life cycle

The **merozoite** recognizes a **specific receptor site** on the **RBC** and exposes several organelles for attachment. The red cells becomes **deformed** and the merozoite enters through an invagination of the RBC membrane

For *P. vivax*, the receptor site on the RBC is associated with the *Duffy Blood Group Antigen*. **Over 90% of Africans are Duffy Negative** and are **resistant** to Vivax Malaria

Pathogenicity and Virulence

Pathological process is the result of **erythrocytic cycle**

1. Hemolysis

RBC invasion by merozoites induces changes in **cytoskeleton** and decreases in **deformability**

Pathogenicity and Virulence (cont)

Increased capillary permeability, which allows fluid to lead into surrounding tissues, causing congestion in blood vessels and **tissue infarction and necrosis**

2. Liberation of Metabolites

Rupture of RBCs bring on **malarial paroxysm** and **increased hemosiderin** in the **liver, spleen**, and other organs, which become **black** in color



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Pathogenicity and Virulence (cont)

Lysis of cells & phagocytosis of cell remnants, debris, and hemozoin congest the **kupffer cells** of the liver, resulting in **hepatosplenomegaly**. Tears of splenic capsule may occur due to trauma or coughing & may require surgery

3. Immunologic Response

Antigens of *P. falciparum* induces **cytokines**, which stimulate the release of **TNF** or **Cachexin**, which causes **fever, paroxysms, HA, pains, and prostration**

Pathogenicity and Virulence (cont)

TNF produces **neurologic symptoms**, which play a role in **cerebral malaria**

Repeated attacks of malaria produce **anemia**

Immune complexes are formed and may be deposited in the **kidney**

4. Formation of Malarial Pigment

In infected RBC, **Hgb** is **digested** forming the pigment **hemozoin**, which can get deposited to various organs

Hemozoin formation **depletes iron stores**, which leads to **anemia**

VIRULENCE FACTORS

Knobs develop for **adhesion** to parasitized and non-parasitized cells and platelets

Pathogenicity and Virulence (cont)

P. falciparum **erythrocyte membrane Pr1 (PfEMP1)** is the **most adhesive protein** among the knobs

Knobs contain **Histidine-Rich Pr (HRP)**

In *Falciparum*, **cytoadherence** causes plugging of **small vessels** by masses of parasitized RBCs and RBCs sticking to **endothelium** causing: **ischemia of the brain, heart, kidney, lungs, and GIT**

PREFERENCE

Vivax and Ovale	Only reticulocytes and young RBC
Malariae	Senescent cells

Pathogenicity and Virulence (cont)

infections with these 3 parasites (vivax, ovale, and malariae) limits number of RBCs that can be parasitized to less than 3% of all erythrocytes

Falciparum and Knowlesi	All ages of RBC
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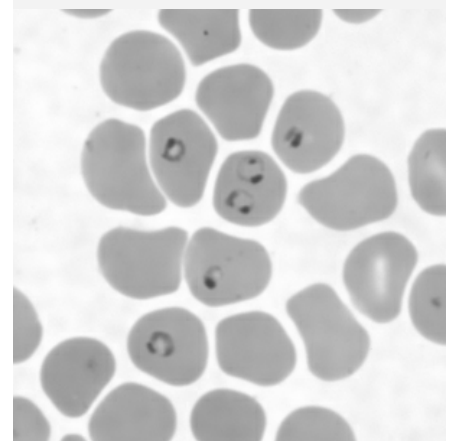
Results in increased parasitemia, early anemia, and increased severity of complications and mortality

Morphology

Ring Form (Early Trophozoite)

Earliest stage after invasion of RBC. The ring has a dot-like nucleus of red chromatin	<i>Falciparum</i> ring: headphone type with a double chromatin dot .
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Plasmodium Falciparum Ring Form



Headphone type is also seen in *P. knowlesi*



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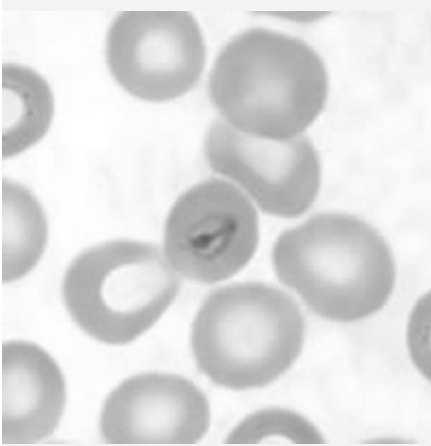
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Plasmodium Malariae Bird's Eye Ring Form



Mature Trophozoites

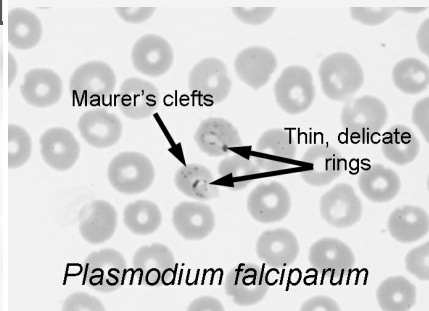
The erythrocyte hemoglobin is metabolized to produce a darkly staining pigment **hemozoin**

Maurer's Clefts: *P. falciparum* infections containing older ring-form trophozoites and asexual stages

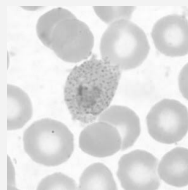
Schüffner's Dots: seen in *P. vivax* and *P. ovale*

Parasite nutrition is hemoglobin and the metabolite is hemozoin or hematin

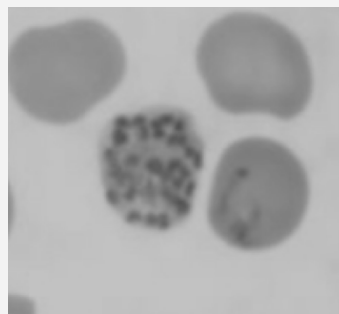
Maurer's Clefts



Schüffner's Dots



Schizont



When parasite divides and shows multiple masses of **nuclear chromatin**. Mature schizonts contain **merozoites**.

Rarely seen in peripheral blood of *P. falciparum* except in severe infections

Falciparum schizonts stay in capillaries of organs and muscles. If schizont **spills into general circulation**, indicates **bad prognosis**

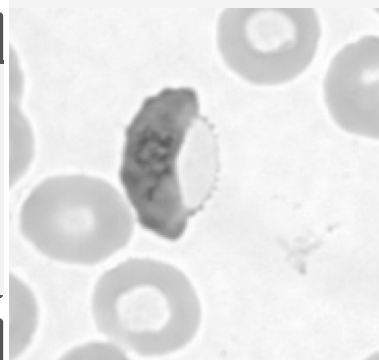
Gametocytes

Sexual form ingested by the mosquito (**infective stage for mosquito**)

Macrogametocyte: nucleus is **dense and compact**

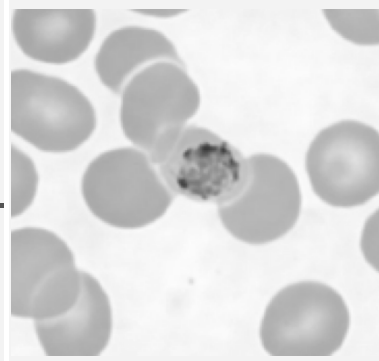
Microgametocyte: nucleus is a **pale, loose network**

Falciparum Gametocyte



Banana shaped

Knowlesi Gametocyte



Gametocytes of **vivax, ovale, malariae**, and **knowlesi** are similar, except that **malariae** is **smaller** and **darker**

Diagnosis

MICROSCOPIC IDENTIFICATION

Gold standard

Specimens can be taken any time

Giemsa stain is preferred for visualization of certain structures (ex. Maurer's Clefts)

If high degree of synchrony exist, late developmental stages can be demonstrated in the repeated smears every 4-6hrs (**not recommended for falciparum**)

Repeated testing is not recommended if parasites are not found initially

Blood smears are obtained to monitor response to treatment

Thick Film:
reveals if
parasite is
present and
is most
efficient
method of
detection

Thin Film:
identifica-
tion of
species

MALARIAL
RDT



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Diagnosis (cont)

Uses Immunochromatography to detect Plasmodium specific antigens in a finger prick blood sample

Antigens utilized are: HRP II, pLDH, Plasmodium Aldolase

HRP II: produced by **trophozoites** and young **gametocytes** of *P. falciparum*

pLDH: produced by both **sexual and asexual** stages, and can distinguish between falciparum and non-falciparum. Sensitive for severe malaria

Plasmodium Aldolase: PA + HRP II = PMA which has 90% specificity and can be performed in 30 minutes

SEROLOGY Cannot differentiate past from current infection

Indirect Hemagglutination (IHA)

Diagnosis (cont)

Indirect Fluorescent Antibody Test (IFAT)

ELISA

PCR low parasitemia or mixed infection

Plasmodium Falciparum

MALIGNANT TERTIAN MALARIA

Also known as subtertian or Aestivoautumnal Malaria

Deadliest species of plasmodium

Causes **50%** of all malaria cases

Erythrocytic cycle of **48hrs**

Found in tropics, subtropics, and sub saharan Africa

Infected cell is **same size** as normal RBC

Associated with the development of **Burkitt's Lymphoma**

RING-FORM TROPHOZOITES

Headphone Type: Rings may possess 1 or 2 chromatin dots

Plasmodium Falciparum (cont)

They may be found on the **periphery** of the RBC (**accolé, appliqué**) and **multiply-infected RBCs** may be seen.

Usually **no enlargement** of infected RBC

Maurer's Clefts: can be seen in infections w/ older ring-form trophozoites, and resembles the **Schüffner's dots** but are **larger** and **coarser**

DEVELOPING AND OLDER TROPHOZOITES

Remain in ring-form, but may become **thicker and more compact**

Amount of **pigment** and **chromatin** may also **increase**

SCHIZONTS

Schizogony **DOES NOT** take place in the **peripheral blood**, but in **capillaries** of **organs** and **muscles**.

The only stages seen in peripheral blood are **rings** and **gametocytes**

Contain anywhere from **8-24 merozoites**

Plasmodium Falciparum (cont)

Mature schizont usually fills 2/3 of infected RBC

GAMETOCYTES

Crescent or **sausage** or **banana** shaped

Pigment is **more coarse and concentrated** in **macrogametocyte** than microgametocyte

Laveran's Bib: remnants of host RBC

Plasmodium Ovale

OVALE MALARIA

Has an erythrocytic cycle of **48hrs**

Has a **hypnozoite** stage in the liver, which is the cause of relapse

Infected RBC is **slightly enlarged** compared to normal RBC

Found in Tropical Africa, West Africa, South America, and Asia

RING-FORM TROPHOZOITES

Contains **single chromatin dot**. Difficult to differentiate from vivax.



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Plasmodium Ovale (cont)

Multiply-Infected RBC may be seen

MATURE TROPHOZOITES

As trophozoite matures, infected RBCs may exhibit **fimbriation** and **Schüffner's dots**

SCHIZONTS

Similar to *P. vivax*, though tend to contain fewer merozoites (**4-16, on average 8**)

Elongation to an **oval shape** and **fimbriation** are common

Schüffner's dots can be observed with proper staining

GAMETOCYTES

Difficult to distinguish from *P. vivax*

Slight enlargement of infected RBC

Plasmodium Vivax

BENIGN TERTIAN MALARIA

Erythrocytic cycle of **48 hours**

Plasmodium Vivax (cont)

Hypnozoite: persistent dormant stage in the liver, which is the cause of **relapse**

Predominant malarial species

Infected **RBC** is **enlarged** due to affinity for **young RBCs or reticulocytes**

RING-FORM TROPHOZOITES

Has a **thick cytoplasm w/ single, large chromatin dot**. Difficult to distinguish from *P. ovale*

Cytoplasm becomes **ameboid** and **Schüffner's dots** may appear as trophozoites mature

LATE TROPHOZOITES

Developing trophozoites become **amoeboid**, with **pseudopodial processes** and **large vacuoles**. Schüffner's dots are visible with proper staining

band-form appearance of trophozoite may occur

May be mistaken for *P. Malariae* or *P. knowlesi*

Plasmodium Vivax (cont)

SCHIZONTS

Pigment is organized in **1-2 clumps**

Mature schizonts contain **12-24 merozoites**, each of which contains a **dot of chromatin** and a **mass of cytoplasm**

GAMETOCYTE

Macrogametocyte: **round to oval** and usually fill host cell. Infected RBC is larger and **cytoplasm** is usually a **darker blue** with **fine brown pigments** throughout

Microgametocyte: usually the size of an **uninfected RBC** and has a **paler blue, pink or grey cytoplasm**

Plasmodium Knowlesi

KNOWLESI MALARIA

Vector is ***A. balabacensis* & *A. maculatus***

Primate malaria in macaques in South East Asia

Infects **ALL** stages of **RBC**, and causes **severe malaria**

Plasmodium Knowlesi (cont)

Erythrocytic cycle has **quotidian** pattern (every **24hrs**)

Non-relapsing due to absence of hypnozoites

Microscopically **indistinguishable** from *P. malariae* (Use PCR to distinguish)

Quotidian pattern results in **high levels of parasitemia** and **severe disease with fatal consequence**

TREATMENT

Chloroquine

Primaquine

Should be treated aggressively and urgently

RING-FORM TROPHOZOITES

Normal to 0.75x smaller than uninfected RBC

Rings may show **double chromatin dots**

Appliqué forms may be seen as well as rectangular rings harboring one or more **accessory chromatin dots**

RBC may be **Multiply-infected**



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Plasmodium Knowlesi (cont)	Plasmodium Malariae (cont)	Plasmodium Malariae (cont)	Clinical Manifestations (cont)
OLDER, DEVELOPING TROPHOZOITES	Old cells are preferentially infected**	Mature schizonts nearly fill the normal-sized host	INCUBATION PERIOD
Band forms may appear similar to <i>P. malariae</i>	RING-FORM TROPHOZOITES	GAMETOCYTES	time between sporozoite injection and appearance of clinical symptoms (8-40)
Sinton and Mulligan's stippling may appear	Have 1 chromatin dot and a cytoplasm ring that is thicker than <i>P. falciparum</i> .	Compact and <i>fills*</i> host RBC	falciparum: 8-15 days
SCHIZONTS	Infected RBC is normal to smaller	Sometimes, there is a reduction in size of the infected RBC	vivax: 12-20 days
Sinton and Mulligan's stippling may appear	Bird's-eye forms may appear	Cytoplasm stains blue and the chromatin is pink to red .	ovale: 11-16 days
10-16 merozoites	MATURE TROPHOZOITES	Abundant dark pigment may be scattered throughout the cytoplasm	malariae: 18-40 days
Gametocytes	Rounded chromatin and compact cytoplasm	Band forms are present in <i>malariae</i> , <i>vivax</i> , and <i>knowlesi</i>	knowlesi: 5 days- few weeks
Mature Macrogametocyte: usually spherical and fill the host RBC. Cytoplasm stains blue and eccentric nucleus stains red	As the trophozoite matures, the cytoplasm may elongate across the host RBC, forming a band-form	Clinical Manifestations	PRODROMAL SYMPTOMS
Microgametocyte: smaller and cytoplasm stains pale pink , while nucleus stains a darker red	Pigment granules become larger and tend to have a more peripheral arrangement	PRE-- PATENT PERIOD	Weakness, exhaustion, aching bones, limbs, and back; loss of appetite; nausea; vomiting
Plasmodium Malariae	Zeimann's stippling may be present	Interval from sporozoite injection to detection of parasites in blood	malaise, backache, diarrhea, and epigastric discomfort
QUARTAN MALARIA	SCHIZONTS	<i>P.falciparum</i> : 11-14 days	MALARIAL PAROXYSM
Erythrocytic cycle of 72hrs	6-12 merozoites , often arranged in a rosette or irregular cluster	vivax: 11-15 days	1. Cold Stage
Found in subtropical and temperate areas		ovale: 14-26 days	Sudden feelings of coldness and apprehension
Infected cells are normal to smaller in size than most RBCs		malariae: 3-4 weeks	
		knowlesi: 9-12 days	



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Clinical Manifestations (cont)

Mild shivering quickly turns to violent teeth chattering and shaking of entire body

Vomiting may occur

Rigors last for 15min-1hr, then stops

2. Hot Stage or Flush Phase

Patient becomes hot (41° C), with headache, palpitations, tachypnea, epigastric discomfort, thirst, nausea, and vomiting

Skin is hot and flushed, patient is confused and delirious

Lasts 2-6hrs

3. Sweating Stage

Defervescence and diaphoresis or profuse sweating occurs

Clinical Manifestations (cont)

Temperature lowers in the next 2-4hrs and symptoms diminish accordingly

Total duration is 8-12hrs

Periodicity of attack only occurs if patient is left untreated

Interval length is determined by length of erythrocytic cycle

falciparum, vivax and ovale – 48 hours

malariae – 72 hours

knowlesi – 24 hours

COMPLICATIONS

Vivax, ovale and quartan malaria are relatively **benign**

Knowlesi

Severe thrombocytopenia, jaundice, deranged liver enzymes.

Clinical Manifestations (cont)

Acute Respiratory Distress Syndrome with tachypnea, hypoxemia, and pulmonary infiltrates on CXR.

Acute Renal Failure with elevated serum creatinine

Hypotension

Acidosis

Chronic Malariae

immune-complex deposition on the glomerular walls, leading to **nephrotic syndrome** in children

Falciparum

Cerebral Malaria (requires prompt administration of **quinidine IV** and then **quinine PO**)

Anemia

Acute Renal Failure (tubular necrosis and nephrotic syndrome)

Clinical Manifestations (cont)

Blackwater fever: from massive intravascular hemolysis and hemoglobinuria

Dysenteric Malaria

Algid Malaria: rapid development of hypotension and impairment of vascular perfusion

Pulmonary Edema

Tropical Splenomegaly Syndrome

Hyperparasitemia (>10-20% of RBC mean high mortality rate)

Hypoglycemia

Treatment

Proper use of the antimalarial drugs is based on knowledge of their effects on the parasite at **various stages of the life cycle**



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Treatment (cont)	Treatment (cont)	Treatment (cont)	Treatment (cont)
<p>Supportive therapy: destroy the parasites as they enter the bloodstream; effective against the erythrocytic stages.</p> <p>Clinical Cure: full and rapid elimination of plasmodium from the blood to prevent complications</p> <p>Radical cure: elimination of not only the bloodstream infection but the tissue stages in the liver as well.</p>	<p>1. Quinine blood schizonticide against all five species of human malarial parasites. Side effects of treatment include tinnitus and headache, vertigo</p> <p>2. Quinidine</p> <p>3. Chloroquine</p> <p>4. Amodiaquine</p> <p>5. Mefloquine effective against both chloroquine-sensitive and – resistant strains of <i>P. falciparum</i> and <i>P. vivax</i>. It is also effective against <i>P. malariae</i> and <i>P. ovale</i></p> <p>6. Doxycycline</p> <p>7. Proguanil Prevents development of oocysts in mosquito</p> <p>8. Halofantrine</p>	<p>9. Artemisinin effective against <i>P. falciparum</i>, <i>P. knowlesi</i> and <i>P. vivax</i>, and in patients with cerebral malaria</p> <p>TISSUE SCHIZO-NTICIDES destroy the developmental stages in the liver</p> <p>1. Primaquine effective against the hypnozoites of <i>P. vivax</i> and <i>P. ovale</i>.</p> <p>*GAMETOCYCIDES</p> <p>1. Primaquine gametocytocidal for all five species of malaria parasites and acts to render the patient noninfectious to the mosquito</p> <p>UNCOMPLICATED P. FALCIPARUM</p> <p>1. Atovaquone-Proguanil (Malarone™) 4 adult tabs po qd x 3 days</p>	<p>2. (Coartem™) 1 tablet = 20mg artemether and 120 mg lumefantrine . 4 tabs initial dose, followed by 4 tabs as second dose 8 hours later, then 4 tabs po bid for the following 2 days.</p> <p>3. plus one of the following: Doxycycline, Tetracycline, or Clindamycin (for pregnant)</p> <p>4. Mefloquine</p> <p>CHLOROQUINE-SENSITIVE FALCIPARUM OR UNCOMPLICATED P. MALARIAE</p> <p>1. Chloroquine phosphate</p> <p>UNCOMPLICATED P. VIVAX OR OVALE</p> <p>1. Chloroquine phosphate plus Primaquine phosphate</p> <p>CHLOROQUINE RESISTANT VIVAX</p> <p>1. Quinine sulfate plus either Doxycycline or Tetracycline plus Primaquine phosphate</p>



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Treatment (cont)

SEVERE MALARIA

1. Quinidine gluconate plus one of the following: Doxycycline, Tetracycline, or Clindamycin
2. Exchange transfusion has been recommended for very severe falciparum malaria associated with high parasitemia (> 10% of RBCs)

P. KNOWLESII

Uncomplicated

1. Chloroquine
2. Primaquine

Severe

1. Quinine
2. Artemether-Lumefantrine

MALARIA PROPHYLAXIS

1. Atovaquone/Proguanil
2. Doxycycline
3. Chloroquine
4. Mefloquine
5. Primaquine



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