The Pigment of the Malaria Parasite *Plasmodium berghei*

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SUMMARY: The pigment of *Plasmodium berghei*, a recently discovered parasite of a wild rodent, *Thamnomys surdaster*, in the Belgian Congo, has been isolated from the infected red cells of young rats in which the strain is maintained in the laboratory. It has been shown by chemical and spectroscopical evidence to be haematin.

Early work by Brown (1911) indicated that the pigment occurring in the spleen and liver of malarial patients infected with *Plasmodium falciparum*, the parasite of malignant tertian malaria, was haematin. Sinton & Ghosh (1934a, b) and Ghosh & Sinton (1934) extracted the pigment from the red cells of monkeys heavily parasitized with *P. knowlesi* and established that this substance was closely similar to or identical with haematin. Devine & Fulton (1941) and Morrison & Anderson (1942) presented further data in support of the nature of this monkey pigment. From spectroscopic and other evidence, that present in *P. gallinaceum* of chickens was also shown by Devine & Fulton (1942) to be haematin. Because the above authors used dilute alkali for extraction, which is known to bring about changes in the properties of this substance, Rimington & Fulton (1947) reinvestigated the pigment of *P. knowlesi* and of *P. gallinaceum* using concentrated phenol for extraction, and established beyond reasonable doubt that the vinyl side chains in the haematin remained unaffected. Recently a plasmodium, which parasitizes the wild rodent *Thamnomys surdaster* in the Congo, has been described by Vinecke & Lips (1948) and named by them *Plasmodium berghei*. It readily infects mice, rats, hamsters and other laboratory animals, and has proved useful in chemotherapeutic experiments. This note describes a chemical investigation of the pigment present within the parasite.

**Experimental**

Hooded rats of weight 60–80 g. were inoculated intraperitoneally with infected blood from a donor rat and 1 week later, when the infection had reached a peak, the animals were anaesthetized and then bled by cardiac puncture in presence of heparin. The infected and non-infected red cells obtained by centrifugation were laked in distilled water and repeatedly washed till free from haemoglobin. The residue was kept under ethanol till sufficient was available for extraction of pigment. This extraction was carried out as described by Rimington & Fulton (1947), using a concentrated solution of phenol prepared by adding 10 ml. water to 90 g. of crystalline phenol. The pigment-containing material was mixed at room temperature with twice its
volume of the 90% phenol solution, and was frequently shaken during a period of several hours. The process of extraction was repeated till the absorption band at 627 mμ was faint or absent. The combined phenol extracts were then mixed with twice their volume of ethanol and filtered through a no. 4 sintered glass funnel to remove precipitated protein. The solution was then dialysed against tap water in a cellophan bag and the contents, which separate into two layers, were frequently mixed. Some stringy material appeared at this time but contained only small amounts of pigment and was not further investigated. Dialysis was continued against distilled water till turbidity occurred in the coloured layer or till its volume was markedly decreased. On mixing this coloured layer with a large volume of water a fine precipitate was produced. The latter was collected by centrifugation and washed repeatedly in distilled water and finally dried in vacuo. It was noteworthy that the washed red cells containing *P. berghei* were very much paler than a corresponding volume of material containing *P. knowlesi* or *P. gallinaceum* and furnished much less pigment than the last two samples. Difficulty is frequently experienced in observing the pigment of some strains of *P. berghei* in stained smears of blood, but in that at present employed it was more abundant and as stated by the original authors "on peut distinguer dans le cytoplasme des grains très fins d'un pigment noir". In gametocytes it occurs in smaller grains and is more widely scattered as shown in Pl. 1. The photomicrographs of the malarial parasites were recorded on Barnet Orthochromatic plates, employing a Leitz 2 mm. 1.4 N.A. oil immersion apochromatic objective, and a Leitz projection ocular. A Wratten no. 45 filter was used to obtain maximum contrast.

**Chemical examination of the residue after dialysis**

A small quantity of the residue was dissolved in 0.01 N-NaOH and one-fifth of its volume of pyridine was added followed by a little sodium dithionite. The spectrum of pyridine haemochromogen appeared (band 557 mμ). The remainder of the residue was subjected to the Grinstein (1947) procedure for the isolation of protoporphyrin ester from haemoproteins. The crystals obtained had m.p. 219° (uncorr.) and absorption maxima in chloroform 631.2, 576.7, 540.4, 503.9 mμ and appeared to be identical in all respects with protoporphyrin IX dimethyl ester. The original pigment is thus identified as haematin.

**Comment**

The amount, appearance and location of pigment in different species of malaria parasites and in different developmental forms of the same parasite show wide variations. The effect on the host cell also differs. In human infections the enlarged pale cell in *P. vivax* infections in which fine yellowish brown pigment is formed contrasts with the normal appearance of the red cell and the presence of abundant coarse dark pigment in infections with *P. malariae*. Because of the apparent anomaly in the case of the latter parasite regarding the appearance of the red cell and the amount of pigment formed, some authors have found it difficult to accept that the pigment arises from the
J. D. Fulton & C. Rimington—Malaria pigment. Plate 1
Malaria pigment

haemoglobin of the host cell. In some strains of *P. berghei* the pigment is frequently difficult to detect; the original authors described it as dark and in very fine granules. In certain monkey and fowl parasites the pigment is characteristic. In all malaria parasites so far examined the material has been shown to be haematin.

Acknowledgement is made to Dr J. E. Falk for help in isolation of the protoporphyrin, and to Mr M. R. Young for the photomicrographs.

REFERENCES


EXPLANATION OF PLATE

Fig. 1. Schizont of *P. berghei* from hamster bone marrow showing deeply stained chromatin and large granules of lighter pigment scattered throughout the cytoplasm. \(\times 2250\).

Fig. 2. Schizont of *P. berghei* from rat blood showing finer pigment granules than that from hamster. \(\times 2250\).

Fig. 3. Female gametocyte of *P. berghei* from hamster blood showing abundant fine pigment granules and small peripheral nucleus. Three other infected cells present. \(\times 2250\).

Fig. 4. Male gametocyte from rat blood with abundant pigment granules throughout the cytoplasm and large deeply stained nucleus. A doubly infected red cell is shown below it. \(\times 2250\).

(Received 17 July 1952)