The Association of Clinical Pathologists: 91st general meeting

The 91st general meeting was held at Imperial College, London, from 19 to 21 September 1973. Abstracts of the scientific communications follow. The Presidential Address, 'Haematology—trends and opportunities', given by Professor E. K. Blackburn, and the Dyke Foundation Lecture, 'Men and machines', given by Professor N. H. Martin, are published in the November issue.

Giant Nuclear Mas ses in the Lungs and Blood in Malignant Malaria

H. B. GOODALL (Royal Infirmary, Dundee) Numerous large naked nuclear masses have been found in the lungs of two patients dead from malignant (subtietian, falciparum) malaria. These structures are probably derived from vascular endothelial cells damaged and shed in the process of disseminated intravascular coagulation which is the pathogenetic mechanism causing the complications of malignant malaria, particularly the cerebral damage. Ten buffy coat smear and 30 thick films of blood from one of these patients taken during the terminal illness were all contain nuclear masses. microscopy of such simple preparations, or of quantitative modifications of them, or of more elaborate cell concentrates, appears to offer a new method for the detection of disseminated intravascular coagulation in malaria, a method to supplement current diagnostic tests, eg, the estimation of fibrin degradation products.

Clinical and Diagnostic Importance of Synovial Fluid Examination

B. VERNON-ROBERTS (The London Hospital Medical College, London) The examination of a sample of synovial fluid from a joint effusion can establish or exclude the diagnosis of crystal synovitis or septic arthritis; it can often provide sufficient evidence for a definite diagnosis in effusions of uncertain aetiology; and it can indicate the degree of disease activity in rheumatoid arthritis and other inflammatory arthropathies.

In the normal knee joint there is less than 3 ml of clear, pale yellow, highly viscous fluid. It contains less than 750 leucocytes/mm³. The majority of cells are small lymphocytes, and the remainder are neutrophil polymorphs and mononuclear phagocytes.

In osteoarthritis the fluid is usually increased in amount, but remains clear and often retains its high viscosity. The total and differential cell count seldom differs from the normal.

In rheumatoid arthritis the fluid is increased in amount, is cloudy due to increased cellularity, and has a reduced viscosity so that it is often watery in consistency. Clot may form due to its increased content of fibrinogen. The total cell count is usually raised above 3000 leucocytes/mm³, and may exceed 50 000 leucocytes/mm³. Neutrophil polymorphs comprise over 50% of the cells present in fluids having high cell counts associated with active arthritis, but may comprise less than 50% of the cells in fluids having low cell counts during inactive phases of the disease. The content of albumin, globulin, IgA, IgG, IgM, and complement in the fluid is increased.

During acute episodes of crystal synovitis the fluid may be purulent in appearance, but the total and differential cell count is largely dependent upon the number of crystals present. Crystals of sodium urate (gout) and calcium pyrophosphate dihydrate (pseudogout) are identified by their specific optical properties.

In septic arthritis the total cell count may be quite low in the early stages and thus may easily be confused with an inflammatory arthropathy or crystal arthritis. In all doubtful cases, the presence of microorganisms is investigated using appropriate special stains and by culture.

In posttraumatic effusions red blood cells may only be numerous for a short time after the traumatic episode has occurred. Red blood cells may also be numerous due to the aspiration procedure, acute synovitis of any cause, haemophilia, or villonodular synovitis. Breaching of the articular surfaces following trauma or bone necrosis may be revealed by the presence of numerous intracellular fat droplets following the uptake of bone marrow fat by synovial fluid leucocytes.

Platelet Aggregation with Ristocetin

JANE HUGHES, ELAINE WILSON, T. W. BARROWCLIFFE, AND PENELOE STANEFORTH (The Royal Free Hospital, London) Following the reports of Howard and Firkin (1971, 1973) that Ristocetin induced platelet aggregation was a valuable diagnostic test for von Willebrand's disease (vWD), 30 previously diagnosed von Willebrand patients were reinvestigated. Standard investigations of blood coagulation and platelet function were carried out, as well as measurement of the factor VIII-related protein and Ristocetin-induced aggregation. With a final Ristocetin concentration of 1·0 mg/ml, aggregation of von Willebrand platelet-rich plasma (PRP) was reduced in all patients (less than 10% aggregation compared with the control range of 30-70%). In 29 out of 30 patients the findings supported the hypothesis that the von Willebrand factor is in the plasma, since Ristocetin-induced aggregation of von Willebrand platelets suspended in normal (N) platelet poor plasma (PPP) was normal, while aggregation of N-platelets in von Willebrand PPP was abnormal. The exception to this was patient 19, in whom mixing experiments indicated a platelet defect (see table).

Twenty-eight patients had reduced or absent levels of factor VIII related protein, as reported by Zimmerman, Ratnoff, and Powell (1971).

Since the two patients with normal levels of VIII-related protein, cases 17 and 19, had hereditary patterns consistent with an autosomal dominant inheritance, they were dissimilar from the cases of von Willebrand's disease with