# Clinical review

#### Recent advances in haematology

Drew Provan, Denise F O'Shaughnessy

Haematology is a diverse specialty embracing clinical and laboratory aspects of adult and paediatric disease, both malignant and non-malignant. We describe here some of the progress that has been made in diagnostic and therapeutic strategies. Molecular advances are continuing at a phenomenal rate (exceeding the rate of progress in therapeutics), offering highly sensitive methods for disease detection and, in some disorders, prospects of our ethnough gene theraps.

#### Methods

We used information from recent key meetings, including those of the British Society for Elaematology and the American Society of Hermatology; leading articles in major harmatology journals; and discussion with colleagues. The choice of topics covered is largely personal, and osting to space restrictions we have not included every "advance" within the specialty.

#### Amaginalias

Distinguishing between anaemia due to iron deficiency and anaemia of chronic disease is a difficult but common problem in medical practice. Anaemia of chronic disease is complex and involves inflammatory extokines," reduced marrow response to erythropoietin, reduced red cell life span, and impaired reuse of iron.1 In typical, uncomplicated iron deficiency anaemia the haemoglobin concentration, mean cell volume, and serum ferritin and iron concentrations are reduced with raised total iron binding capacity; unforturnately cases are often not typical, and the results of these tests may seem conflicting. Furthermore, if a patient with amarmia of chronic disease is also iron deficient, the parameters for diagnosing iron deficiency are altered, making the diagnosis difficult; often a borse marrow aspirate, stained for iron, is the only method for accurately assessing from status. This is expensive, time consuming, and unpleasant for the patient. Recently the serum transferrin receptor assay has been developed, enabling more accurate assessment of iron status in this group of patients,

Serum ferritin, a 480 kDa multisubunit protein, represents the body's iron storage pool. A reduced serum ferritin concentration generally indicates depletion of the iron stores Ferritin, however, is an "acute phase protein," whose concentration is raised in inflammatory disorders, in a patient with, for example, active rheumatoid disease the ferritin concentration may be normal (or even raised) even if the patient is

#### Recent advances

from deficiency arusemia can more be distinguished from arusemia of chronic disease by using serum transferrin receptor assays

A recently discovered human herpesvirus may have a role in development of raveloma

Additional thrombotic risk factors—for example, factor V and prothrombin gene mutations, hyperhomocysteinaemia—have been identified

Newly designed DNA probe arrays have the potential to allow rapid automated detection of mutations

Antisense therapy, ribosyme technology, and other novel treatment methods are being developed

truly iron deficient. The transferrin receptor assay aims to distinguish clearly between simple iron deficiency and anaemia of chronic disease in most cases.

How does the transferrin receptor away work? Transferrin is a protein that carries plasma iron. Cellsthat require iron express the transferrin receptor on the cell surface." The receptor binds plasma transferrin with bound iron atoms, after which the transferrin receptortransferrin complex is internalised (fig 1). After unloading the iron atoms, the transferrin receptor-transferrin complex moves back to the cell surface and dissociates, leaving transferrin free to pick up further iron atoms, In iron deficiency anaemia the number of transferrin receptors increases (is upregulated),' whereas in anacmia of chronic disease the number of transferring receptors remains normal. Assays for transferrin receptors may be performed on serum with an enzyme linked irruntinosorficut assoc (ELISA), allowing large numbers of samples to be batch tested. According to available data, the test seems as reliable as testing bone marrow assistate for diagnosing from deficiency and should be valuable in potients with riscurnatology disorders and others with inflammatory disorders.

#### Haemophilia

Haemophilias A (factor VIII deficiency) and B (factor IX deficiency, Christmas disease) are X linked blooding Department of Hamanicky, Scattlange, Scattlange, Scattlange, Scattlange, Scattlange, SCO SLV Dere Provint, since Intensit, since Intensit, since Intensit, SCO SLV Draw Provint CISIonal States (CISIonal States), see a since SCO SCATTLANGE, SCATTLANG

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**Brendan G. Carr** 

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Nathan and Oski's Hematology of Infancy and Childhood E-Book Stuart H. Orkin, David G. Nathan, David Ginsburg, A. Thomas Look, David E. Fisher, Samuel Lux, 2008-12-16 To address the exponential growth in the fields of pediatric hematology and oncology this classic reference has been separated into two distinct volumes With this volume devoted strictly to pediatric hematology and another to pediatric oncology you ll keep you on the cutting edge of these two specialties The completely revised 7th edition of Nathan and Oski s Hematology of Infancy and Childhood is now in full color and provides

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**Hematology** E. Cacciola, A. B. Deisseroth, R. Giustolisi, 1994-01-13 **Hematology E-Book** Ronald Hoffman, Edward J. Benz, Leslie E. Silberstein, Helen Heslop, Jeffrey Weitz, Mohamed E. Salama, 2022-10-27 Extensively revised comprehensive content from leading global contributors ensures that Hematology 8th Edition remains your 1 choice for expert guidance in all areas of this rapidly advancing subspecialty This edition reflects the numerous advances that are redefining the field and dramatically influencing new approaches to diagnosis treatment and outcomes Well illustrated and clinically focused it details the basic science and clinical practice of hematology and hematopoietic cellular therapy covering virtually all aspects of hematology in one definitive resource Covers all hematologic disorders including comprehensive discussions of hematologic malignancies individualized patient care cell based therapies transplantation transfusion medicine hemostasis thrombosis and consultative hematology in one convenient volume Provides state of the art guidance from global experts at the forefront of the latest research and clinical practice Provides extensive updates throughout on basic science research advances in molecular diagnostics new drugs immunotherapies personalized medicine laboratory medicine transfusion medicine stem cell transplantation and clinical treatment for all hematologic malignancies and non malignancies Contains new chapters on gene editing the impact of mitochondria on hematopoiesis myelodysplastic syndrome myeloproliferative neoplasm overlap syndromes immunotherapy and management of its toxicities transfusion medicine in sickle cell disease principles of radiation therapy and COVID 19 including complications of vaccination and its impact on the hematologic system Discusses many new advances in the field including details and the future of gene therapy for hemophilia gene editing for sickle cell disease and thalassemia the evolution of cellular therapy use of cells transfusion medicine vs protein therapy gene sequencing immunotherapy and new targeted drugs Includes more decision making algorithms for formulating diagnoses and personalized treatment plans for those highly complex disorders that require individualized approaches Addresses the effects of aging on hematopoiesis and on the manifestations of a variety of hematologic disorders Discusses cardio oncology and its impact on the treatment of patients with hematologic disorders Presents relevant basic science as background for clinical application in later sections An eBook version is included with purchase The eBook allows you to access all of the text figures and references with the ability to search customize your content make notes and highlights and have content read aloud Wintrobe's Clinical Hematology John P. Greer, 2009 Now available in a thoroughly revised Twelfth Edition Wintrobe's Clinical Hematology continues to be an industry leader with its ability to correlate basic science with the clinical practice of hematology With the first edition of Wintrobe's Clinical Hematology published in 1942 clearly establishing hematology as a distinct subspecialty of Internal Medicine the latest edition continues the influence of the Wintrobe name and helps to set this book apart from the competition With its strong focus on the clinical aspects of hematology the book has generated a strong following among internists and general practitioners who want a single resource to consult for their patients who present any blood related disorder The Twelfth Edition is in full color for the first time

boasts a new editorial team and includes expanded coverage of new medications and four new chapters on Newborn Anemias Pathology of LHC Spleen Tumors and Myeloproliferative Disorders and Mast Cell Disease A companion Website will offer the fully searchable text and an image bank Hematology Marshall A. Lichtman, Laurence A. Boxer, Edward Henderson, Sanford J. Shattil, Jerry L. Spivak, 2000-04-26 This book is a centennial volume celebrating the enormous progress made in hematology in the 20th century It is edited by Marshall Lichtman a distinguished senior hematologist past president of the American Society of Hematology and co editor of the leading text in the field Hematology is a compendium with commentaries of the most important papers published in the field from 1900 1999 The book will be useful for reference many of the older papers can no longer be found in most libraries yet are still referred to in current publications especially review articles as well as teaching The Editor and a team of associate editors have included the most important papers covering eight categories anemia phagocytic cells platelets coagulation and thrombosis lymphocytes and immune disorders transfusion medicine hematologic malignancies and therapeutics and laboratory developments Each paper is accompanied by a 1 2 page commentary explaining its impact and references to the developments that resulted Key Features Contains 86 landmark articles from the last 100 years of research in clinical hematology Includes expert commentaries discussing the impact of each article Cites approximately 1000 preceding or subsequent articles of consequence in the commentaries Includes the English translations of nine articles originally published in other languages Provides easy access to several Transplantation in Hematology and Oncology T. Büchner, H. Jürgens, W.E. papers that may no longer be found in libraries Berdel, J. van de Loo, J. Ritter, J. Kienast, J. Vormoor, 2012-12-06 Some three decades after bone marrow transplantation was introduced in the field of hematology and oncology transplantation today continues to rapidly grow and expand into a variety of new modalities Peripheral blood has been established as an effective source of autologous progenitor cells Furthermore the graft versus leukemia effect has resulted in novel strategies of adoptive immunotherapy for cancer Finally approaches to gene transfer and therapy are utilizing transplantation methodologies and can augment their effects Current results new developments and perspectives are presented in this volume Conventional and innovative experimental approaches the past and the future of bone marrow transplantation are reviewed and discussed by leading representatives **Archives of** Pathology, 1928 Research Grants Index National Institutes of Health (U.S.). Division of Research Grants, 1970

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