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ILLUSTRATED REVIEW



Illustrated State-of-the-Art Capsules of the ISTH 2021 Congress

Sriram Krishnaswamy PhD¹ | Walter Ageno MD² | Yaseen Arabi MD³ \checkmark | Tiziano Barbui MD⁴ | Suzanne Cannegieter MD, PhD⁵ \checkmark | Marc Carrier MD, MSc, FRCPC⁶ | Audrey C. Cleuren PhD⁷ | Peter Collins⁸ | Laurence Panicot-Dubois⁹ \checkmark | Jane E. Freedman MD¹⁰ \checkmark | Kathleen Freson PhD¹¹ \checkmark | Philip Hogg PhD¹² | Andra H. James MD¹³ \checkmark | Colin A. Kretz PhD¹⁴ \checkmark | Michelle Lavin MB, PhD, FRCPath^{15,16} \checkmark | Frank W. G. Leebeek MD, PhD¹⁷ | Weikai Li PhD¹⁸ \checkmark | Coen Maas PhD¹⁹ | Kellie Machlus PhD²⁰ \checkmark | Michael Makris MD²¹ \checkmark | Ida Martinelli MD, PhD²² | Leonid Medved PhD²³ | Marguerite Neerman-Arbez PhD²⁴ | James S. O'Donnell PhD^{25,26,27} | Jamie O'Sullivan PhD²⁸ \checkmark | Madhvi Rajpurkar MD^{29,30} | Verena Schroeder PhD³¹ | Paul Clinton Spiegel Jr PhD³² \checkmark | Simon J. Stanworth MD^{33,34,35} | Laura Green MD^{36,37} \checkmark | Anetta Undas MD, PhD³⁸

¹Hematology, Department of Pediatrics, Children's Hospital of Philadelphia, Perelman School of Medicine, University of Pennsylvania, Philadelphia, PA, USA ²University of Insubria, Varese, Italy

³King Abdulaziz Medical City, Ministry of NGHA, King Saud Bin Abdulaziz University for Health Sciences, Riyadh, Saudi Arabia

⁴Research Foundation, Papa Giovanni XXIII Hospital, Bergamo, Italy

⁵Depertments of Clinical Epidemiology and Thrombosis & Haemostasis, Leiden University Medical Center, Leiden, The Netherlands

⁶Department of Medicine, Ottawa Hospital Research Institute, University of Ottawa, Ottawa, ON, Canada

⁷Life Sciences Institute, University of Michigan, Ann Arbor, MI, USA

⁸School of Medicine Cardiff University, Haemophilia Centre, University Hospital of Wales, Cardiff, UK

⁹Faculty of Pharmace, AMU C2VN INSERM 1263, Marseille, France

¹⁰Vanderbilt University Medical Center, The Albert Sherman Center, Worcester, MA, USA

¹¹Center for Molecular and Vascular Biology, KU Leuven, Leuven, Belgium

¹²Charles Perkins Centre, University of Sydney, Sydney, NSW, Australia

¹³Duke University, Durham, NC, USA

¹⁴TaARI, McMaster University, Hamilton, ON, Canada

¹⁵National Coagulation Centre, St. James's Hospital, Dublin, Ireland

¹⁶Irish Centre for Vascular Biology, RCSI, Dublin, Ireland

¹⁷Department of Hematology, Erasmus MC, University Medical Center, Rotterdam, The Netherlands

¹⁸Washington University in St. Louis Medical School, St. Louis, MO, USA

¹⁹University Medical Center Utrecht, Utrecht, The Netherlands

²⁰Vascular Biology Program and Harvard Medical School, Boston Children's Hospital, Boston, MA, USA

²¹University of Sheffield, Sheffield, UK

²²Hemophilia and Thrombosis Center, IRCCS Fondazione Ca' Granda Ospedale Maggiore Policlinico, Milano, Italy

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²³Center for Vascular and Inflammatory Diseases and Department of Biochemistry and Molecular Biology, University of Maryland School of Medicine, Baltimore, MD, USA

²⁴Deartment of Genetic Medicine and Development, Faculty of Medicine, University of Geneva, Geneva, Switzerland

²⁵Haemostasis Research Group, Irish Centre for Vascular Biology, School of Pharmacy and Biomolecular Sciences, Royal College of Surgeons in Ireland, Dublin, Ireland

²⁶National Children's Research Centre, Our Lady's Children's Hospital, Dublin, Ireland

²⁷National Centre for Coagulation Disorders, St James's Hospital, Dublin, Ireland

²⁸Irish Centre for Vascular Biology, School of Pharmacy and Biomolecular Science, Royal College of Surgeons in Ireland, Dublin, Ireland

²⁹Children's Hospital of Michigan, Central Michigan University, Detroit, MI, USA

³⁰Wayne State University, Detroit, MI, USA

³¹Department for BioMedical Research, University of Bern, Bern, Switzerland

³²Western Washington University, Bellingham, WA, USA

³³Transfusion Medicine, NHS Blood and Transplant, Oxford, UK

³⁴Department of Haematology, Oxford University Hospitals NHS Foundation Trust, Oxford, UK

³⁵Radcliffe Department of Medicine, NIHR Oxford Biomedical Research Centre, University of Oxford, Oxford, UK

³⁶Transfusion Medicine, NHS Blood and Transplant (London) and Barts Health NHS Trust, London, UK

³⁷Blizzard Institute, Queen Mary University of London, London, UK

³⁸Jagiellonian University Medical College, Krakow, Poland

Abstract

This year's Congress of the International Society of Thrombosis and Haemostasis (ISTH) was hosted virtually from Philadelphia July 17-21, 2021. The conference, now held annually, highlighted cutting-edge advances in basic, population and clinical sciences of relevance to the Society. Despite being held virtually, the 2021 congress was of the same scope and quality as an annual meeting held in person. An added feature of the program is that talks streamed at the designated times will then be available online for asynchronous viewing. The program included 77 State of the Art (SOA) talks, thematically grouped in 28 sessions, given by internationally recognized leaders in the field. The SOA speakers were invited to prepare brief illustrated reviews of their talks that were peer reviewed and are included in this article. The topics, across the main scientific themes of the congress, include Arterial Thromboembolism, Coagulation and Natural Anticoagulants, COVID-19 and Coagulation, Diagnostics and Omics, Fibrinogen, Fibrinolysis and Proteolysis, Hemophilia and Rare Bleeding Disorders, Hemostasis in Cancer, Inflammation and Immunity, Pediatrics, Platelet Disorders, von Willebrand Disease and Thrombotic Angiopathies, Platelets and Megakaryocytes, Vascular Biology, Venous Thromboembolism and Women's Health. These illustrated capsules highlight the major scientific advances with potential to impact clinical practice. Readers are invited to take advantage of the excellent educational resource provided by these illustrated capsules. They are also encouraged to use the image in social media to draw attention to the high quality and impact of the science presented at the congress.

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Splanchnic vein thrombosis

Walter Ageno MD

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Anticoagulant Therapy for Splanchnic Vein Thrombosis: Summary of Suggested Treatment Strategies

(Adapted from Di Nisio et al JTH 2020)



For references, see Di Nisio et al.¹

Yaseen Arabi MD



Prevention of venous thromboembolism in critically ill patience involves multi-modality approach. A systematic review of randomized controlled trials (RCTs) demonstrated that early mobilization in critically ill patients was associated with a significant reduction in deep vein thrombosis (DVT) (RR 0.16, 95% CI 0.06, 0.47).² Avoidance of unnecessary use of central venous catheters, especially in the femoral vein, is an important aspect of DVT preventive strategy. One RCT found that subclavian compared to femoral venous catheterization in critically ill patients was associated with a significant reduction in catheter-related thrombosis documented by ultrasonographic examination (RR 0.09, 95% CI 0.02, 0.36).³ Pharmacologic thromboprophylaxis with unfractionated or low-molecular-weight heparin, compared to no thromboprophylaxis, also reduces the incidence of DVT (RR 0.51, 95% CI 0.41, 0.63).⁴ Data on the effectiveness of pneumatic review that included observational studies demonstrated lower DVT with intermittent pneumatic compression (RR 0.34, 95% CI 0.19, 0.60).⁵



Myeloproliferative neoplasm-associated thrombosis

Tiziano Barbui



PV, polycythemia vera; ET, Essential Thrombocythemia; PMF, primary myelofibrosis; MPN, myeloproliferative neoplasms Pts/yr, patients/year

For references, see Barbui et al,⁶ Tefferi et al,⁷ and De Stefano et al.⁸

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Venous thrombo-embolism; risk in premenopausal women: Questions still unanswered

Suzanne Cannegieter MD PhD



VTE: Venous thrombo-embolism; PCOS: Polycystic ovary syndrome For references, see Samuelsson et al,⁹ Scheres et al,¹⁰ and Roach et al.¹¹

Primary thromboprophylaxis: Who, what, and how?

Marc Carrier MD, MSc, FRCPC



Venous thromboembolism (VTE) is associated with significant morbidity, mortality and healthcare utilization among ambulatory patients with cancer initiating systemic chemotherapy. Direct oral anticoagulants and low molecular weight heparins have been shown to be safe and effective to prevent cancer-associated thrombosis in this patient population.¹² Hence, the use of primary thromboprophylaxis should be considered to decrease the risk of VTE and tailored to minimize the risk of bleeding.¹² Patients should be educated about signs and symptoms of cancer-associated thrombosis and stratified according to their underlying risk of VTE and bleeding for potential consideration of primary thromboprophylaxis implementation in ambulatory patients with cancer initiating systemic chemotherapy.¹³

DOAC: Direct oral anticoagulants; KS: Khorana Score; LMWH: Low molecular weight heparin; VTE: Venous thromboembolism.

Molecular analysis of vascular gene expression

Audrey C. Cleuren



Vascular heterogeneity, particularly in the endothelial cell (EC) compartment, has long been recognized yet difficult to study given the poor accessibility of these cells. (A) The Ribotag mouse model¹⁴ enables evaluation of gene expression profiles directly *in vivo* in a cell-specific manner via translating ribosome affinity purification (TRAP). (B) EC-TRAP combined with high-throughput RNA sequencing provides an accurate *in vivo* snapshot of organ-specific endothelial gene expression programs. In addition to the extensive EC heterogeneity across organs under physiologic conditions shown here, our data also showed vascular bed-specific EC reactivity after lipopolysaccharide-induced endotoxemia (ref¹⁵).

Hemostatic management of postpartum hemorrhage

Peter Collins

Hemostatic management of postpartum hemorrhage

Peter Collins MD



Postpartum hemorrhage (PPH) is caused by obstetric complications but may be exacerbated by impaired hemostasis. Hypofibrinogenemia is the commonest coagulopathy associated with PPH and occurs early in abruptions and amniotic fluid embolism. Depletion of other coagulation factors, thrombin generation and platelets is uncommon until large bleeds have occurred.¹⁶

Early hypofibrinogenemia predicts progression to severe hemorrhage, however, laboratory Clauss fibrinogen is usually too slow to be clinically useful during rapid bleeding. Point-of-care viscoelastometric hemostatic assays (VHA) allow surrogate measurement of fibrinogen and predict severe outcomes.¹⁷ A double blind RCT showed that fibrinogen >2 g/L is adequate for hemostasis during PPH.¹⁸

An all Wales quality improvement programme involving 60,000 deliveries combined accurate measurement of blood loss with VHAs to guide early fibrinogen replacement. It was associated with reduced massive PPH (\geq 2500 ml) by 23%, decreased red cell and FFP transfusion by 22% and 58%, respectively and reduced severe anemia (Hb <80 g/L) by 33%.

Mechanistic insights from 3-D visualization of thrombus formation in cancer

Laurence Panicot-Dubois



For references, see Carminita et al¹⁹ and Palacios-Acedo et al.²⁰

Platelets: Influenza and other viral responses

Jane E. Freedman, MD

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Growing mechanistic and clinical data demonstrate that platelets perform various immune functions during infection. A platelet can form heterotypic aggregates with various types of immune cells including monocytes, neutrophils, eosinophils, and dendritic cells. Platelets participate in innate and adaptive immunity and act as immune cells during viral infections. Platelets may internalize ssRNA viruses including influenza, HIV, severe acute respiratory syndrome coronavirus-2 (SARS-CoV-2) and encephalomyocarditis. For some viruses, this internalization leads to lysosomal degradation of the viral coat and activation of the pathogen-associated molecular pattern receptor, TLR7 (Toll-like receptor 7). Activation of TLR7 by influenza also results in C3 (complement 3) release from platelets that leads to complement cascade activation and release of neutrophil DNA which can contribute to aggregates.

Hemostatic phenotypes and genetic disorders

Kathleen Freson

2021: Genetic landscape of Inherited Platelet Disorders



More than 60 genes have already been discovered as cause of an inherited platelet disorder (IPD).²¹ Many of these genes are widely expressed and are associated with broader clinical symptoms than causing solely a bleeding tendency. In this presentation examples will be discussed on how next generation sequencing has proven successful with its implementation in clinical diagnostics^{22,23} and gene discovery.²³ Still at least half of the IPD patients receive no genetic diagnosis. Ideas will be put forward on how to tackle the challenges ahead that include discoveries in the noncoding genome space and setting up improved disease models for IPD that will allow (automated) deep phenotyping.

Functional disulfide bonds in hemostasis and thrombosis

Philip Hogg PhD



For references, see Butera and Hogg,²⁴ Hogg²⁵

Disparities in pregnancy outcomes: Differences by condition and community

Andra H. James MD



Blood donations per 1000 population by countries' income: a measure of blood availability



Both the maternal and fetal outcomes of pregnancy vary according to a pregnant woman's community and her condition. The most devastating outcome is the death of a month. On 2017, there were approximately 295,000 maternal deaths with dramatic differences in maternal mortality based on the region of the world, the country, and women's underlying conditions.²⁶ Worldwide, the leading cause of maternal death is hemorrhage. Ninety-nine percent of maternal deaths and 99% of those due to hemorrhage occur in low- or middle-income countries. Whether a hemorrhage originates from inside the uterus (80%), from laceration or incisions (20%), or from an underlying coagulopathy (less than 1%), and acute acquired coagulopathy will evolve unless the hemorrhage is not available, because besides the usual obstetric measures, blood, hemostatic medication and hematologic expertise are necessary to save mothers' lives.^{27,28}

Specificity of ADAMTS13 and regulation of ADAMTS13 function

Colin A. Kretz



Pregnancy, postpartum and periods: Current challenges in the management of women with Von Willebrand disease

Michelle Lavin MB, PhD, FRCPath



Challenges in the management of women with VWD

Women are disproportionately impacted by Von Willebrand disease (VWD) due to gynaecological bleeding. Heavy menstrual bleeding (HMB) is the most frequently reported and highest scoring bleeding symptom for women with VWD yet optimal treatment strategies remain uncertain.²⁹ In pregnancy, there remains controversy regarding the ideal therapeutic plasma Von Willebrand factor (VWF) target at delivery. While thresholds similar to surgery are often utilized, this approach fails to account for the physiological pregnancy-induced increase in plasma Von Willebrand factor (VWF) levels, with median plasma VWF levels >200–250 IU/dl in healthy women at delivery.³⁰ The limitations of current approaches for women with VWD are reflected in primary postpartum haemorrhage (PPH) rates, which remain increased even when replacement therapy is used.³¹

Postpartum, as plasma VWF levels return, to baseline women with VWD are at a markedly increased risk of secondary PPH following discharge. As women with VWD may be normalized to HMB, recognition of secondary PPH may be delayed or missed.

The impact of aging and inflammation on plasma Von Willebrand factor levels

Frank W.G. Leebeek



In the general population Von Willebrand factor (VWF) levels rise with aging, especially above the age of 40.³² As is shown in the figure, several mechanisms have been suggested to be responsible for this increase. This may be increased release of VWF from the endothelium, or decreased VWF clearance. Mechanisms contributing to increase of VWF are endothelial dysfunction, comorbidities (hypertension, diabetes), weight gain, atherosclerosis and inflammation.^{32,33} This age-related increase is also observed in patients with type 1 Von Willebrand disease (VWD) and may result in (near) normal levels in elderly VWD patients.³⁴ It is still disputed whether this rise is associated with attenuation of the bleeding tendency. Results of an observational study on bleeding symptoms over one year have shown that bleeding was not reduced in type 1 VWD patients above the age of 65 compared to those <65 years, however prospective data are still lacking.³⁴

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Structural basis of antagonizing the vitamin K catalytic cycle for anticoagulation

Weikai Li PhD



The catalytic cycle of vitamin K epoxide reductase (VKOR) and inhibition by vitamin K antagonists are accompanied with redox-state and conformation changes. The hemisphere (pink) and cylinder (grey) illustate VKOR luminal and transmembrane domain, respectively. VKOR reduces vitamin K epoxide (KO) to quinone (K) and then to hydroquinone (KH2). A catalytic cysteine forms covalent and charge-transfer complex with KO and K, generating mercapto adducts of 3-hydroxyl K (S-KOH) and K (S-KH), respectively. Their binding induces a closed conformation that juxtaposes all cysteines (SH or S-S) for unimpeded electron transfer to reduce the mercapto adduct. VKOR becomes fully oxidized and changes to an open conformation that releases the reaction product. Warfarin competes with the substrates for the partially-oxidized enzyme. Unlike the substrates, warfarin binds also to the fully-oxidized enzyme and removes it from the enzyme pool. The bound warfarin locks HsVKOR in both redox states into a closed conformation.³⁵⁻³⁷

Pathogenic factor XII mutations: Form determines dysfunction

Coen Maas





Kellie Machlus PhD



During inflammation, platelets are activated and release extracellular vesicles (EVs). These EVs enter the bone marrow (BM), where they bind to resident BM cells including megakaryocytes (MKs). These plasma-originating EVs help communicate changes in the circulation directly into the BM, and may contribute to BM reprogramming during inflammation.

Current and future haemophilia treatment options: Clinician perspective

Michael Makris MD

HEMOPHILIA TREATMENTS



Cerebral venous sinus thrombosis

Ida Martinelli MD, PhD



Abbreviations: CVST, cerebral venous sinus thrombosis; CT, computed tomography; MRI, magnetic resonance imaging

Fibrin(ogen)-endothelial cell interactions in inflammation

Leonid Medved, PhD



Two of the proposed fibrin(ogen)-dependent pathways of leukocyte transmigration are based on the bridging mechanism. The first one suggests that fibrinogen bridges leukocytes to the endothelium through the interaction with endothelial receptor ICAM-1 and leukocyte integrin Mac-1 to promote leukocyte transmigration⁴¹ (left panel). The second one suggests that fibrin degradation product E_1 fragment promotes leukocyte transmigration by bridging leukocytes to the endothelium through the interaction with endothelial VE-cadherin and leukocyte integrin CD11c (left panel), and the β 15-42 fragment inhibits this interaction and thereby inflammation.⁴² Our studies revealed that interaction of fibrin with the endothelial VLDL receptor promotes transendothelial migration of leukocytes through the fibrin-VLDL receptor-dependent pathway (left panel), identified two monoclonal antibodies inhibiting this interaction, and clarified the molecular mechanism underlying this pathway and the inhibitory role of β 15-42 in this pathway⁴³ (right panel). These antibodies exhibited significant anti-inflammatory properties and may represent potential therapeutics for treatment of fibrin-dependent inflammation.

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Fibrin(ogen) in human disease

Marguerite Neerman-Arbez PhD

Fibrin(ogen)	Mechanism	Consequence	Outcome
	Mutations	Quantitative defects: - low or absent fibrinogen Qualitative defects: - abnormal clot structure Amyloidosis	Bleeding, Thrombosis Bleeding, Thrombosis Renal disease, hypertension, CVD
	Inappropriate post-translational modifications	Abnormal clot structure	Bleeding, Thrombosis
	High levels (e.g. inflammation)	Hypercoagulability, metabolic dysfunction	CVD Cancer: worse outcome
	Abnormal localisation - in CNS - In kidney	Inflammation, demyelination Amyloidosis	Neurological disease Renal disease
	Interaction with virulence factors	Limits antimicrobial role of fibrinogen	Microbial infections
	Extra-hepatic expression (carcinoma)	Increases cancer growth and invasive potential	Cancer: worse outcome

With fibrin, produced by thrombin-mediated cleavage, fibrinogen plays important roles in many physiological processes.^{44,45} Formation of a stable blood clot, containing polymerised and cross-linked fibrin, is crucial to prevent blood loss and drive wound healing upon injury. Balance between clotting and fibrinolysis is essential. Several diseases are the consequence of altered levels of fibrinogen, others are related to structural properties of the molecule. Inflammation leads to elevated circulating levels of fibrinogen and hypercoagulability, a risk factor for cardiovascular disease (CVD). The source and localisation of fibrin(ogen) also has clinical implications. Fibrin(ogen) has been associated with cancer development and progression. While fibrin(ogen) is implicated in defense against pathogens, in other settings it enhances bacterial virulence.

Von Willebrand factor modulates adhesion of malaria-infected erythrocytes to endothelial cells

James S. O' Donnell



VWF strings modulate adhesion of malaria-infected erythrocytes

Endothelial cell monolayer

Markedly elevated plasma VWF:Ag levels and VWF propeptide levels are present in children with severe *Plasmodium falciparum* malaria, consistent with acute endothelial cell (EC) activation and Weibel Palade body (WPB) secretion.⁴⁶ Higher VWF levels correlate with worse clinical outcomes. Pathological ultra-large (UL-) VWF multimers are also a feature of cerebral malaria.⁴⁷ In vitro studies have demonstrated that UL-VWF strings on the surface of activated EC can recruit platelets. Subsequently, the platelet-decorated UL-VWF strings can then tether malaria-infected red blood cells (IRBC) under physiological shear stress.⁴⁸ In particular, *P. falciparum* erythrocyte membrane protein-1 (PfEMP-1) on the IRBC interacts with platelet CD36. Collectively, these findings support the hypothesis that VWF plays role in regulating microvascular sequestration of IRBC in children with cerebral malaria.

Image created with BioRender.com.

Von Willebrand Factor structure-function in the regulation of cancer metastasis

Jamie O'Sullivan



For reference, see Patmore et al⁴⁹

Pulmonary embolism in children

Madhvi Rajpurkar MD

Pulmonary Embolism in Children Current knowledge and future perspectives Madhvi Rajpurkar, MD



For references, see Biss et al,⁵⁰ Carpenter et al,⁵¹ Rajpurkar et al⁵²

Intermolecular interactions that stabilize multimeric FXIII

Verena Schroeder PhD

Intermolecular Interactions that Stabilize Multimeric FXIII



For references, see Handrkova et al,⁵³ Li et al,⁵⁴ Schroeder et al⁵⁵

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Factor VIII structure: Determinants of inhibitor development

Paul Clinton Spiegel, Jr. PhD



Hemophilia A is an X-linked bleeding disorder that affects 1 in 5,000 males worldwide due to a deficiency in blood coagulation factor VIII (fVIII), an essential protein for the proteolytic activation of factor X to Xa. Through the past 20 years, the structure and function of fVIII has come into focus. New atomic-level structural findings illustrate a detailed domain organization of fVIII structure,⁵⁶ novel localized and domain-scale conformational changes, the molecular nature of the fVIII/Von Willebrand factor complex, and pathogenic antibody epitopes.^{57,58} Further structural characterization of fVIII circulatory complexes will uncover the fundamental basis for its procoagulant cofactor function and may aid in next-generation bioengineering efforts to improve fVIII stability and circulatory half-life while minimizing its immunogenicity. These efforts may prove vital for both fVIII replacement and gene therapy approaches.

Do platelet transfusions work?

S.J. Stanworth and Laura Green



A number of randomised trials have evaluated the risk-benefit ratio for platelet transfusion for prophylaxis and treatment of major bleeding. Trials in some settings have indicated evidence of harm with more 'liberal' use of platelets,⁵⁹ although a benefit to improve outcomes has also been found in major bleeding.⁶⁰ Our understanding of donor, storage and processing characteristics on outcomes following platelet transfusion have been inadequately investigated.⁶¹ We need a better understanding of how platelet transfusions affect both haemostasis and inflammation in patients, to indicate which patients really require platelet transfusions.

Prothrombotic fibrin clot properties and vascular diseases

Anetta Undas MD, PhD



Clinical conditions associated with prothrombotic fibrin clot phenotype

Fibrin clot structure characterized by fiber diameter and pore size differs between healthy persons and patients with thromboembolic diseases. Prothrombotic fibrin clot phenotype is associated with faster formation of denser fibrin mesh, relatively resistant to lysis, as reflected by prolonged clot lysis time (CLT). Increased plasma fibrin clot density has been reported in patients with prior or acute thromboembolic events.

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WA has received grant funding from Bayer and payments from Bayer, Portola, Aspen, Sanofi, BMS/Pfizer, Daiichi Sankyo; MC has received grant funding from Leo Pharma, Pfizer, BMS and consulting fees from Leo Pharma, Pfizer, Bayer, BMS, Servier, Sanofi; ACC has received funding from the American Heart Association; PC has received grant funding from CSL Behring, Werfen (TEM International), Haemonectics, the Welsh Government and personal fees from CSL Behring, Werfen; JEF has received funding from the American Heart Association; KF has received funding from Sobi (Swedish Orphan Biovitrum); ML has received consulting fees from Sobi, Tremeau Pharmaceuticals, Takeda; FWGL has received grant funding from CSL Behring, Takeda/Baxalta, uniQure, SOBI and consulting fees from CSL Behring, Biomarin, Takeda, UniQure; KM has received funding from the American Society of Hematology, NIH and consulting fees from STRM.BIO; MM has received grants from Bayer, BPL, CSL Behring, Kedrion, NovoNordisk, Octapharma, Pfizer, Roche, Sobi, Takeda and consulting fees from Novo Nordisk, Grifols, Sanofi; LM has received funding from the NIH; JO has received grant funding from LEO Pharma; VS has received funding from the Swiss National Science Foundation; all other authors declare no conflicts of interest.

TWITTER

Yaseen Arabi (wysseenarabiya) Suzanne Cannegieter (ws_cannegieter) Laurence Panicot-Dubois (wglaneFreedmanMD) Kathleen Freson (wglaneFreedmanMD) Kathleen Freson (wglandrajames031) Colin A. Kretz (wglandrajames031) Colin

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