Editors’ Choice in the 60th Anniversary Year of Thrombosis and Haemostasis: Past, Present and Future

Christian Weber1  Gregory Y. H. Lip2

1Institute for Cardiovascular Prevention, Ludwig-Maximilians-University, Munich, Germany
2Haemostasis Thrombosis and Vascular Biology Unit, City Hospital, Birmingham, West Midland, United Kingdom


2017 marks a very special year for Thrombosis and Haemostasis: it marked the 60th anniversary of the journal. To commemorate this event, selected section editors were invited to create a modern portrait of the journal, reflecting the milestones of the past and encouraging a glance into future across the many topics of the journal’s scope. The main directions reflected in the anniversary editorial1 are recapitulated in this year’s Editors’ Choice selection of papers of high interest or citation.

We traditionally start the New Year with various position and consensus papers and current state-of-the-art developments, revealing valuable insights for researchers and clinical specialists. For instance, the role of microvesicles in the pathophysiology of thrombosis and cardiovascular disease has been intensely investigated in light of their value as potential biomarkers. We are therefore delighted to publish a relevant position paper of the European Society of Cardiology Working Group on Atherosclerosis and Vascular Biology, which comprehensively summarizes the current mechanistic knowledge on the formation, composition and function of microvesicles with an endothelial, platelet, red blood cell and leukocyte origin.2

In addition, we also published an executive summary of a joint consensus document from the European Heart Rhythm Association and European Society of Cardiology Working Group on Thrombosis, endorsed by the ESC Working Group on Valvular Heart Disease, Cardiac Arrhythmia Society of Southern Africa (CASSA), Heart Rhythm Society (HRS), Asia Pacific Heart Rhythm Society (APHRS), South African Heart (SA Heart) Association and Sociedad Latinoamericana de Estimulación Cardiaca y Electrofisiología (SOLEACE), addressing the topic of antithrombotic therapy in atrial fibrillation (AF) associated with valvular heart disease.3 This document recommends that the term ‘valvular AF’ is outdated and proposes a new functional Evaluated Heartvalves, Rheumatic or Artificial (EHRA) categorization in relation to the type of oral anticoagulant use in patients with AF, as follows: (1) EHRA Type 1 valvular heart disease which refers to AF patients with ‘valvular heart disease needing therapy with a vitamin K antagonist (VKA)’ and (2) EHRA Type 2 valvular heart disease, which refers to AF patients with ‘valvular heart disease needing therapy with a VKA or a non-VKA oral anticoagulant (NOAC),’ also taking into consideration score risk factors.

New Developments in Anticoagulants

Anticoagulants are the mainstay for the prevention and treatment of thrombosis. In their joint paper, Weitz and Harenberg highlighted the advances of oral anticoagulant therapy, referring to safety issues and reviewing some of the ongoing trials with NOACs, and provided insights into recent research on factors XII and XI inhibitors as potentially novel anticoagulants.4 Poterucha et al in their review focused on heparin interference in the inflammatory pathophysiology of thrombosis development and tended to advancing treatments based on heparin derivatives.5 To complement this, Schulman et al reviewed published as well as ongoing observational studies with NOACs in VTE and discussed limitations in analysis and interpretation.6

With the NOACs, the quest for reversal agents continues, and there was much interest in the paper by Ansell et al7 on ciraparantag, a new ‘universal antidote’ and small molecule that potentially reverses the anticoagulant effects of dabigatran and factor Xa inhibitors.

However, despite the developments with NOACs, the VKAs are not ‘dead’ and we published interesting papers on the pharmacology of tecarfarin, a novel VKA drug.8,9 This complements the clinical data from the EmbraceAC trial.10
Venous Thromboembolism

Venous thromboembolism (VTE) as the third most frequent acute cardiovascular syndrome is associated with a considerable disease burden. In the anniversary issue, Schulman et al recalled therapeutic developments focusing on prophylaxis, attending to new drugs and technical advances in imaging, as well as hospital accreditation and funding based on evidence-based practice.11

Additional analyses from the large trials included a pooled analysis of RE-COVER and RE-COVER II trials comparing dabigatran versus warfarin for acute venous thromboembolism in elderly or impaired renal function patients.12

Apart from clinical trials, real-world observational data on NOACs are increasingly evident. As one notable example, Coleman et al assessed the effectiveness and safety of rivaroxaban versus warfarin in VTE patients in routine practice and came to the conclusion that it more effectively reduces patients’ hazard of both recurrent VTE and major bleeding.13

Trujillo-Santos et al investigated the use of NOACs under real-life conditions, including nonrecommended doses or regimens and unsealed the influence on VTE occurrence and subsequent outcomes.14 We also published the design paper for RE-COVERY DVT/PE, a prospective observational study of acute venous thromboembolism with a focus on dabigatran etexilate.15

Metabolic-Vascular Disease

Endothelial cell metabolism has emerged as an important regulator of angiogenesis. In their review, Breier et al provided insights into angiogenesis during obesity-related metabolic dysfunction and into signaling pathways linking cell metabolism to endothelial function.16 A set of publications highlighted the impact of hepatic alterations on thrombosis. Ambrosino et al figured out the importance of specific VTE prevention strategies for cirrhotic subjects,17 whereas an interesting contribution to elucidating the molecular and pathogenic mechanisms underlying nonalcoholic steatohepatitis (NASH) was provided by Geys et al, demonstrating that ADAMTS13 deficiency in obese mice induces hepatic microthrombosis.18,19 Lallukka et al demonstrated that obesity and insulin resistance rather than liver fat increases circulating coagulation factor activities.20 Finally, adipose tissue inflammation starts with cell accumulation. Having disclosed a role of leukocyte integrin Mac-1 in macrophage accumulation in adipose tissue in a murine model, Wolf et al21 remained sceptical concerning the net effect of integrin blockade in cardiometabolic disease.

Stroke Prevention in Atrial Fibrillation

The evolution of stroke and bleeding risk assessment over the past decades was reviewed by Lip et al,22 culminating in a practical management pathway to help streamline and simplify decision making for stroke prevention in patients with AF with three simple steps and to aid decision making for stroke prevention in AF (referred to as the Birmingham ‘3-step’).

Biomarker scores represent an attractive prognostic tool for bleeding risk assessment. The performance of the new ABC-bleeding score was validated and compared with the HAS-BLED score in a ‘real-world’ setting of anticoagulated AF patients with long-term follow-up, which concluded that HAS-BLED performed significantly better in predicting major bleeding.23 In the largest ‘real-world’ study on apixaban effectiveness and safety to date, Li et al showed that apixaban significantly attenuates stroke risk and major bleeding compared with warfarin independent from risk subgroups and dose regimens.24

Therapeutic Strategies for Atherosclerosis and Atherothrombosis

Atherosclerosis and atherothrombosis continue to be the leading causes of death worldwide. In their review, Weber et al provided historical overview, spotted the range of currently available state-of-the-art therapies and highlighted the promising therapeutic strategies aimed at reduction of the residual risk that still persists despite current therapeutic options. Jamasbi et al underscored their expectations for novel antiplatelet drugs selectively inhibiting arterial thrombosis without interfering with normal haemostasis.25

The nature of atherosclerosis had been extensively studied through sophisticated transgenic animal models. As an example, Winkels et al revealed the new atherosclerosis-modulating properties of CD70 through altering macrophage function.26 Pointing out the recent failures in translating various anti-inflammatory therapeutic strategies for use in humans demands to retain a healthy scepticism regarding the inflammatory causality underlying human atherosclerosis despite the recent success of the CANTOS trial, Santovito and Weber advised to keep in mind that ‘things may not always be what they appear’.28

Antithrombotic Therapy for Acute Coronary Syndrome

Plaque erosions and ruptures accompany arterial thrombus formation in the coronary arteries resulting in acute coronary syndrome. Sibbing et al highlighted a discrepancy between current guideline recommendations favouring potent platelet inhibition in ACS and the utilization of the respective drugs in clinical practice and pointed out the development of optimized antiplatelet treatment strategies and their utilization in the real world.29 Standard care for treating non-ST elevation myocardial infarction patients is represented by dual antiplatelet therapy. Interestingly, chewing versus an equal dose of traditional oral administration enhances inhibition of platelet aggregation after administering a ticagrelor.30

Beyond Cardiovascular Disease

The biochemical characterization of the proteolytic pathways that constitute blood coagulation is followed by identifying and validating appropriate targets for improving global health through their application in haemostasis and thrombosis pathologies. In their review, Ten Cate et al evaluated the
concepts providing a modern vision of coagulation, illustrating the importance of the coagulation cascade in cardiovascular pathology through thrombotic as well as atherosclerotic processes and in the response to ischaemia–reperfusion injury. Severe inflammatory complications often determine the disease progression and outcome. The evidence from both basic research and clinical studies highlighted the role of self-extracellular nucleic acids in the crosstalk between immunity and cardiovascular diseases. Preissner et al delivered a portrait of self-extracellular nucleic acids with an emphasis on their role in immune response, inflammation, thrombosis and cardiovascular diseases.

Over the past 60 years, Thrombosis and Haemostasis continues to be a platform for exciting developments in vascular biology and medicine, welcoming next generation of authors with novel and amazing approaches in all aspects of coagulation, haemostatic and vascular research. We look forward to help you in sharing and disseminating your knowledge.

References
19 Wojta J. What ADAMS13 does in the liver... Thromb Haemost 2017;117(01):6