

Novel regulatory mechanism of blood clotting discovered

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GPV modulates thrombin-mediated platelet activation and fibrin formation. On the platelet surface, GPV regulates platelet responsiveness to thrombin by interference with GPIbα-dependent PAR signaling (left). Once GPV is cleaved by thrombin, sGPV dampens thrombin activity and fibrin formation, thereby controlling clot formation (right). Created with BioRender. Credit: *Nature Cardiovascular Research* (2023). DOI: 10.1038/s44161-023-00254-6

Hemostasis is vital to prevent excessive blood loss. However, there is an increased risk for thrombosis if there is an excessive reaction and



uncontrolled formation of fibrin. In a new study published in *Nature Cardiovascular Research*, a research group from Würzburg led by Bernhard Nieswandt have uncovered platelet glycoprotein GPV as a central regulator for hemostasis and thrombus formation, a finding that opens the way to new therapeutic approaches.

When our <u>blood vessels</u> are injured by cuts, abrasions, or bruises, it is vital that the bleeding is stopped, and the wound is sealed. This process is called hemostasis and involves two main components: First, <u>blood</u> platelets attach to the wound edges, form a plug and provisionally seal the injury. Second, blood coagulation or the <u>coagulation cascade</u> is initiated, leading to the formation of long fibrin fibers, which together with platelets seal the wound tightly.

However, if fibrin is formed in excess, e.g., in chronic wounds, thrombosis and subsequent vascular occlusions can occur. Therefore, the strict regulation of fibrin formation is important. How blood clotting is limited was not fully understood until now.

GPV controls thrombin activity and fibrin formation

In this study, the research group led by Professor Bernhard Nieswandt identified a fundamentally new mechanism. "For the first time, we were able to uncover a new switching point that regulates both hemostasis and thrombosis. This switch is glycoprotein V, GPV, which is expressed on the surface of blood platelets. GPV controls the activity of the enzyme thrombin, which is responsible for the formation of fibrin," explains Nieswandt, head of the Institute of Experimental Biomedicine I and member of the board of the Rudolf Virchow Center—Center for Integrative and Translational Bioimaging (RVZ) at the University of Würzburg.

Thrombin is a crucial enzyme in <u>blood clotting</u> and its activity must



therefore be precisely spatiotemporally controlled. Until now, it was known that the surface receptor GPV is cleaved by thrombin during platelet activation. This releases GPV as a soluble form.

However, the physiological function of this receptor was largely unknown. Using genetic and pharmacological approaches, the researchers showed that thrombin-mediated cleavage of GPV limits fibrin formation. By remaining bound to thrombin, soluble GPV alters the activity of thrombin so that it can form less fibrin.

In experimental thrombosis models, soluble GPV was shown to prevent, among other things, the formation of vaso-occlusive thrombi and to lead to a significant protection from experimental stroke and associated brain damage.

Bernhard Nieswandt is convinced that these new findings will change textbook knowledge. He thanks all participating scientists from the RVZ and the University Hospital Würzburg (UKW), who were supported by colleagues from Mainz, Maastricht, and the U.S.

Antibodies against GPV offer great clinical potential in the treatment of disturbed hemostasis

In another approach, the research group generated antibodies against GPV that prevent thrombin-mediated cleavage of GPV. "In our studies, we were able to show that these antibodies increase thrombin activity, resulting in increased fibrin formation. Our idea was therefore to use these antibodies in the context of impaired hemostasis to increase fibrin formation," says Professor David Stegner, head of the Vascular Imaging Group at RVZ and one of the study's last authors.

In addition to genetic causes, a reduced platelet count or impaired



function can also be caused by pharmacological treatment and therefore lead to an impaired hemostasis. Treatment with platelet aggregation inhibitors, such as clopidogrel, which are used to prevent heart attacks or strokes and to treat circulatory disorders, impairs platelet function.

"In an experimental model of hemostasis, our new antibody was indeed able to restore hemostasis under conditions where hemostasis is otherwise not possible. This indicates a support of hemostasis by enhancing thrombin-dependent <u>fibrin</u> formation," adds Dr. Sarah Beck, scientist at the Würzburg Institute for Experimental Biomedicine and first author of the study. "Anti-GPV treatment could have great clinical potential. This will be investigated in more detail in the future."

More information: Sarah Beck et al, Platelet glycoprotein V spatiotemporally controls fibrin formation, *Nature Cardiovascular Research* (2023). DOI: 10.1038/s44161-023-00254-6

Provided by Universitätsklinikum Würzburg

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