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ANGELILLO, Anne, et al.

Abstract

Mechanisms regulating thrombus stabilization remain largely unknown. Here, we report that loss of any 1 of the Gas6 receptors (Gas6-Rs), i.e., Tyro3, Axl, or Mer, or delivery of a soluble extracellular domain of Axl that traps Gas6 protects mice against life-threatening thrombosis. Loss of a Gas6-R does not prevent initial platelet aggregation but impairs subsequent stabilization of platelet aggregates, at least in part by reducing "outside-in" signaling and platelet granule secretion. Gas6, through its receptors, activates PI3K and Akt and stimulates tyrosine phosphorylation of the beta3 integrin, thereby amplifying outside-in signaling via alphaIIbbeta3. Blocking the Gas6-R-alphaIIbbeta3 integrin cross-talk might be a novel approach to the reduction of thrombosis.

Reference


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Role of Gas6 receptors in platelet signaling during thrombus stabilization and implications for antithrombotic therapy

Anne Angelillo-Scherrer,1 Laurent Burnier,1 Nathalie Flores,1 Pierre Savi,2 Maria DeMol,3 Paul Schaeffer,2 Jean-Marc Herbert,2 Greg Lemke,4 Stephen P. Goff,5 Glenn K. Matsushima,6 H. Shelton Earp,7 Christian Vesin,8 Marc F. Hoylaerts,3 Stéphane Plaisance,3 Désiré Collen,3 Edward M. Conway,3 Bernhard Wehrle-Haller,8 and Peter Carmeliet3

1Division of Angiology and Hemostasis, Department of Internal Medicine, Faculty of Medicine and University Hospitals of Geneva, Geneva, Switzerland. 2Department of Cardiovascular Research, Sanofi-Synthélabo Recherche, Toulouse, France. 3Center for Transgene Technology and Gene Therapy, Flanders Interuniversity Institute for Biotechnology, and Center for Molecular and Vascular Biology, University of Leuven, Leuven, Belgium. 4The Salk Institute for Biological Studies, San Diego, California, USA. 5Department of Biochemistry and Molecular Biophysics, Columbia University, New York, New York, USA. 6University of North Carolina Neuroscience Center, and Lineberger Comprehensive Cancer Center, University of North Carolina at Chapel Hill, Chapel Hill, North Carolina, USA. 7Department of Cellular Physiology and Metabolism, University of Geneva Medical School, Geneva, Switzerland.

Mechanisms regulating thrombus stabilization remain largely unknown. Here, we report that loss of any 1 of the Gas6 receptors (Gas6-Rs), i.e., Tyro3, Axl, or Mer, or delivery of a soluble extracellular domain of Axl that traps Gas6 protects mice against life-threatening thrombosis. Loss of a Gas6-R does not prevent initial platelet aggregation but impairs subsequent stabilization of platelet aggregates, at least in part by reducing “outside-in” signaling and platelet granule secretion. Gas6, through its receptors, activates PI3K and Akt and stimulates tyrosine phosphorylation of the β3 integrin, thereby amplifying outside-in signaling via αIIbβ3. Blocking the Gas6–R–αIIbβ3 integrin cross-talk might be a novel approach to the reduction of thrombosis.

Introduction

Formation of a platelet plug initiates hemostasis at sites of vascular injury and triggers pathological thrombosis in ischemic tissue disease. After initial formation of a single platelet monolayer, additional platelets are recruited into the growing hemostatic plug. Without further stabilization, platelet plugs prematurely disaggregate. While platelet plug instability may precipitate rebleeding at sites of vascular injury, inhibition of platelet plug stabilization may be therapeutically attractive to prevent pathological thrombosis. Indeed, agents inhibiting the formation of stable platelet plugs, but leaving behind an intact platelet coverage, might block thrombosis, possibly without eliciting major adverse bleeding effects. The availability of such compounds would be desirable, as currently available antithrombotic agents often cause a bleeding diathesis.

The molecular mechanisms controlling perpetuation of the platelet plug are incompletely characterized. The αIIbβ3 integrin plays a central role in platelet aggregation. In the initial phase, activation of platelets by ADP, thromboxane A2 (TXA2), and thrombin signals “inside-out” via αIIbβ3, thereby increasing its ligand binding and activation (1). In the perpetuation phase, the activated αIIbβ3 transmits critical “outside-in” signals, which commits the platelets to activation, irreversible platelet aggregation, and clot retraction (1). A critical signaling event includes the phosphorylation of tyrosine residues in the cytoplasmic domain of the β3 integrin (2–4). Additional outside-in signaling molecules include the Eph kinases and ephrins, and CD40 ligand (5, 6).

We recently reported that Gas6 enhances the formation of stable platelet macroaggregates in response to various agonists (7). Gas6 binds to the receptor tyrosine kinases Tyro3, Axl, and Mer via its carboxyterminal globular domain (8–10). Each of the Gas6 receptors (Gas6-Rs) has an extracellular ligand-binding domain and a cytoplasmic tyrosine kinase domain, the activity of which is enhanced by Gas6 (9, 11–17). This then leads to further intracellular signaling, including activation of the PI3K and Akt pathways (18–20). The Gas6-R–mediated signaling cascades in platelets remain unknown. Gas6–/– mice are protected against thrombosis but do not suffer from bleeding (7). Mice with a triple deficiency of Tyro3, Axl, and Mer have not been reported to exhibit an abnormal hemostasis under baseline conditions (21), but their response in pathological conditions was not evaluated.

In this study, we determined that Gas6, through its receptors, activates PI3K and Akt and stimulates tyrosine phosphorylation of the β3 integrin, thereby amplifying outside-in signaling via αIIbβ3. Furthermore, we established that a “Gas6 trap” prevents pathological thrombosis, which indicates that this novel cross-talk between the Gas6-Rs and αIIbβ3 integrin may constitute a novel target for antithrombotic therapies.

Results

Loss of a Gas6-R causes rebleeding and protection against thrombosis. Mice lacking any 1 of the Gas6-Rs, i.e., Tyro3–/–, Axl–/–, or Mer–/– mice, did not suffer spontaneous bleeding or thrombosis under baseline conditions. We used 2 bleeding models to investigate the effects of loss of any 1 of the Gas6-Rs on hemostasis. In the first model, we determined the time to cessation of bleeding. When bleeding did not recur within 60 seconds of cessation, bleeding was considered to be arrested. All genotypes had normal bleeding times after tail clipping: 290 ± 56 seconds for WT mice, 300 ± 32 seconds for Tyro3–/– mice, 280 ± 55 seconds for Axl–/– mice, and 290 ± 58 seconds for Mer–/– mice. We subsequently performed a second bleeding model, the tail-clipping model, to compare the time to cessation of bleeding. WT mice bled for 290 ± 56 seconds, while Tyro3–/– mice bled for 300 ± 55 seconds, Axl–/– mice for 290 ± 55 seconds, and Mer–/– mice for 300 ± 57 seconds. Thus, loss of any 1 of the Gas6-Rs, i.e., Tyro3, Axl, or Mer, protected mice against thrombosis.

Nonstandard abbreviations used: Gas6-R, Gas6 receptor; G/l, giga/liter; hAxl-EC-Fc, soluble Axl extracellular domain; PAR, protease-activated receptor; PPP, platelet-poor plasma; PRP, platelet-rich plasma; TXA2, thromboxane A2; αIIbβ3, integrin. Conflict of interest: The authors have declared that no conflict of interest exists.


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were obtained in Tyro3–/– mice by i.v. injection of tissue thromboplastin. Stasis-induced thrombosis was induced by ligation of the inferior vena cava followed by tip transection, the cut end of the tail was immersed in saline at 37°C. Aliquots of saline solution containing blood were sampled every minute for 10 minutes and then every 5 minutes until 30 minutes. The amount of blood loss was determined by measurement of the hemoglobin content of the aliquots of blood collected in saline (mean ± SEM, n = 10). In WT mice, blood loss reached a plateau within 10 minutes, indicating cessation of bleeding. In contrast, Tyro3–/–, Axl–/–, and Mer–/– mice had a tendency to repetitively rebleed after transient hemostasis.

The first bleeding model is not suitable to detect rebleeding after initial hemostasis, we used a second model where the amount of blood loss was estimated every minute for 10 minutes and, thereafter, every 5 minutes until 30 minutes. In WT mice, blood loss increased progressively to reach a plateau within 10 minutes, after which bleeding stopped completely. In contrast, Tyro3–/–, Axl–/–, and Mer–/– mice had a tendency to repetitively rebleed after transient hemostasis. Consequently, Gas6-R–deficient mice lost more blood during 30 minutes after tail clamping than WT mice (95 ± 35 μl in WT mice versus 260 ± 66 μl in Tyro3–/– mice, 260 ± 28 μl in Axl–/– mice, and 270 ± 60 μl in Mer–/– mice; n = 10, P < 0.05). Similar results were obtained in Gas6–/– mice (data not shown).

Two thrombosis models were used to determine the role of Tyro3, Axl, or Mer in pathological thrombosis. In the first model, thrombosis was induced by ligation of the inferior vena cava followed by i.v. injection of tissue thromboplastin. Stasis-induced venous thrombosis results from both coagulation and platelet activation (22). Compared with WT mice, thrombosis in Tyro3–/–, Axl–/–, or Mer–/– mice were on average reduced by more than 90% (n = 10, P < 0.005; Figure 2A). In the second model, platelet-dependent thromboembolism was induced by intrajugular injection of collagen and epinephrine and caused fatal pulmonary embolism within 5 minutes in 80% of WT mice (n = 15). In contrast, fatal pulmonary embolism was observed, after 10–20 minutes, in only 25% of Tyro3–/– mice (n = 15), in 10% of Axl–/– mice (n = 10), and in 20% of Mer–/– mice (n = 12, P < 0.05 versus WT for all Gas6-R–/– mice; Figure 2B). Macroscopic and histological analysis revealed extensive pulmonary thromboembolism in WT mice (Figure 2C). In contrast, no signs of pulmonary embolization were detectable in surviving Axl–/– mice (Figure 2D), or in Tyro3–/– or Mer–/– mice (data not shown). The hemostatic phenotype of Gas6-R–deficient mice was not attributable to genotypic differences in platelet counts, prothrombin and activated partial thromboplastin times, or fibrinogen levels (data not shown).

Although the above model of collagen-induced thromboembolism is recognized to be platelet dependent, we sought to further confirm that the protection against thrombus formation in mice lacking a Gas6-R was predominantly due to altered platelet function. We therefore induced thromboembolism with collagen and epinephrine in Axl–/– mice after platelet depletion and subsequent reconstitution with either WT or Axl–/– platelets (23, 24). Thrombocytopenia was achieved in Axl–/– mice by i.p. injection of 20 mg/kg busulfan on days 1 and 3 and i.v. injection of anti-platelet mAb CV-5H7 on day 3 (25). On day 6, platelet counts dropped to less than 10 giga/liter (G/l). Thromboembolism experiments were performed on day 6 after reconstitution of WT or Axl–/– platelets in Axl–/– thrombocytopenic mice (washed platelets from 3 WT or 3 Axl–/– mice were necessary to reconstitute 1 Axl–/– mouse). Platelet counts in Axl–/– mice after reconstitution with WT platelets or Axl–/– platelets were 546 ± 45 G/l (n = 5) and 583 ± 39 G/l (n = 5), respectively. Fatal pulmonary embolism was observed in 4 of 5 Axl–/– mice reconstituted with WT platelets but, notably, in none of the Axl–/– mice reconstituted with Axl–/– platelets. These results confirm that resistance of Gas6-R–deficient mice to thromboembolism may be largely attributed to a platelet-function defect.

Impaired platelet aggregation and clot retraction in Tyro3–/–, Axl–/–, or Mer–/– mice. Since Gas6 impairs platelet plug formation (7), platelet function of Gas6-R–deficient mice was also studied. Platelets from WT mice dose-dependently aggregated in response to increasing concentrations of ADP, collagen, or the TXA2 analogue U46619, achieving a maximal effect at concentrations similar to those previously reported (7). In contrast, platelets from Tyro3–/–, Axl–/–, or Mer–/– mice failed to irreversibly aggregate in response to low concentrations of ADP (less than 10 μM; Figure 3A), collagen...
(2 μg/ml; Figure 3B), or U46619 (1 μM; Figure 3C), although higher concentrations of these agonists were capable of inducing irreversible aggregation of platelets (Figure 3, A–C). The impaired platelet aggregate formation in Gas6-R–deficient mice resulted from signaling defects most likely upstream of PKC activation or Ca ++ mobilization, since WT, Tyro3−/−, Axl−/−, or Mer−/− platelets aggregated normally in response to PMA or the Ca ++ ionophore A23187 (data not shown). Thrombin (0.01–1.0 IU/ml) and the protease-activated receptor 4–activating (PAR4-activating) peptide AYPGKF (0.25–1.0 mM) (26) comparably stimulated initial platelet aggregation in all genotypes (data not shown). However, clot retraction upon thrombin treatment was significantly impaired in Tyro3−/−, Axl−/−, or Mer−/− mice (Figure 4). These data establish a key role for the Gas6-Rs in the perpetuation of platelet activation and the stabilization of platelet aggregates.

**Normal fibrinogen binding in Tyro3−/−, Axl−/−, or Mer−/− platelets.** Since binding of fibrinogen during initial platelet activation—a process that depends on inside-out signaling to ε130β3 (27)—is critical for platelet aggregate formation, we examined whether the platelet defect of Gas6-R–deficient mice was attributable to impaired fibrinogen binding. However, fibrinogen binding during initial platelet activation was similar in all genotypes. Indeed, platelets of all genotypes bound comparable amounts of Oregon Green 488–conjugated fibrinogen after stimulation with various concentrations of ADP (1.25–20 μM; Figure 5). Thus, loss of a Gas6-R does not seem to affect the initial conversion of ε130β3 to a low-affinity/low-avidity to a high-affinity/high-avidity receptor upon activation of platelets by various agonists.

**Defective outside-in signaling in Tyro3−/−, Axl−/−, or Mer−/− platelets.** To analyze platelet spreading after adhesion to fibrinogen (a process that depends on outside-in signaling via ε130β3), unstimulated platelets were incubated on a fibrinogen-coated surface. Noticeably, fewer platelets from Tyro3−/−, Axl−/−, or Mer−/− mice adhered to fibrinogen, and these failed to spread out as extensively as WT platelets (7). Secretion of ADP from dense granules is essential for the formation of stable macroaggregates after initial formation of small, unstable platelet aggregates. This is accompanied by secretion of adhesion proteins such as fibrinogen, vWFα, and P-selectin from α granules, which further contributes to stable platelet plug formation. Secretion of dense-granule stores (evaluated by measurement of the amount of released ADP (5 μM) or washed human platelets after preincubation with human Axl extracellular domain (hAxl-EC-Fc) or control IgG, revealing that hAxl-EC-Fc reduces platelet aggregation. A representative example of 3 independent experiments is shown. Arrows in A–D indicate addition of the platelet agonists.

**Figure 3**
Effect of the lack of Tyro3, Axl, or Mer on aggregation of WT, Tyro3−/−, Axl−/−, or Mer−/− PRP. (A) Response of WT or Tyro3−/− platelets to 5 μM and 50 μM ADP. (Similar results were obtained with Axl−/− and Mer−/− platelets; data not shown.) (B) Response of WT or Axl−/− platelets to 2 μg/ml and 10 μg/ml collagen. (Similar results were obtained with Tyro3−/− and Mer−/− platelets; data not shown.) (C) Response of WT or Mer−/− platelets to 1 μM and 7 μM TXA2 analogue U46619. (Similar results were obtained with Tyro3−/− and Axl−/− platelets; data not shown.) (D) Aggregation response to ADP (5 μM) of washed human platelets after preincubation with human Axl extracellular domain (hAxl-EC-Fc) or control IgG, revealing that hAxl-EC-Fc reduces platelet aggregation. A representative example of 3 independent experiments is shown. Arrows in A–D indicate addition of the platelet agonists.

**Figure 4**
Effect of the lack of Tyro3, Axl, or Mer on clot retraction. Photographs show the degree of clot retraction after 60, 90, and 120 minutes in WT, Tyro3−/−, Axl−/−, and Mer−/− PRP samples treated with 10 IU/ml thrombin. Tyro3−/−, Axl−/−, and Mer−/− PRP was slower than WT PRP in retraction of clots (n = 3). Clots are surrounded by dotted lines.

Gaskel stimulated tyrosine phosphorylation of the β3 integrin. The above findings that Gaskel-Rs regulate spreading of platelets on fibrinogen and stabilization of the platelet plug suggested that Gaskel-R signaling influenced outside-in signaling by the αIIbβ3 integrin. Upon ligand binding, critical tyrosine residues in the cytoplasmic tail of the β3 integrin become phosphorylated, which then serve as docking sites for downstream signaling molecules. We therefore investigated the effect of Gaskel on β3 integrin tyrosine phosphorylation by immunoblotting using anti-β3 (pY773) phosphospecific antibodies. Since Gaskel, by itself, is unable to promote platelet aggregation (7), treatment of platelets with Gaskel permitted us to study αIIbβ3 tyrosine phosphorylation independently of platelet aggregation. As shown in Figure 9A, treatment of resting platelets with Gaskel stimulated β3 integrin tyrosine phosphorylation in WT platelets. In contrast, such β3 tyrosine phosphorylation did not occur when Tyro3–/– (Figure 9A), or Axl–/– or Mer–/– (data not shown), platelets were treated with Gaskel. The reduced β3 tyrosine phosphorylation in Gaskel-R–deficient platelets was not attributable to a decrease in total β3 integrin expression, as shown by flow cytometry (Figure 9B). Gaskel-R signaling determined the degree of β3 phosphorylation not only in resting platelets, but also in activated platelets after stimulation with thrombin. Indeed, when WT platelets were stimulated with thrombin (which is known to enhance release of Gaskel from the endogenous platelet stores; see ref. 7), the β3 integrin became tyrosine-phosphorylated (Figure 9C). Quantification by densitometry revealed that a lower percentage of the total amount of β3 was tyrosine-phosphorylated (30% ± 4% and 30% ± 4% in WT platelets versus 8% ± 2% and 20% ± 2% in Tyro3–/– platelets after 0.05 and 0.10 U thrombin/ml, respectively; n = 3, P < 0.05). Absence of Gaskel-R significantly reduced but did not, however, abolish tyrosine phosphorylation of the β3 integrin, as higher concentrations and longer times of application of thrombin did stimulate phosphorylation of the β3 integrin (Tyro3–/–, Figure 9C; Axl–/– and Mer–/–, data not shown). Thus, Gaskel-R signaling determines β3 tyrosine phosphorylation and, thereby, also outside-in signaling of αIIbβ3, defects of which result in retarded platelet spreading on fibrinogen.

Gaskel stimulates phosphorylation of its receptors. Since some of the effects of the Gaskel-Rs have been proposed to be ligand-independent and to not require tyrosine phosphorylation of the receptors (28), we analyzed whether Gaskel enhanced the kinase activity of Tyro3, Axl, and Mer. Immunoprecipitation using phosphotyrosine antibodies and subsequent immunoblotting using Gaskel-R–specific antibodies revealed that Gaskel induced tyrosine phosphorylation of Tyro3 (data not shown), Axl (Figure 9A), and Mer (data not shown) in WT platelets. As expected, when a particular Gaskel-R (for instance Axl) was absent, it could not be immunoprecipitated. Remarkably, however, absence of Tyro3 prevented ligand-induced tyrosine phosphorylation of the other Gaskel-Rs (Axl, Figure 9A; and Mer, data not shown). Similarly, when Axl or Mer was absent, Gaskel-induced tyrosine phosphorylation of the residual Gaskel-R failed to occur, suggesting that the Gaskel-Rs cooperate in their activation response (see below).

As this might be due to interdependent expression patterns of each of the Gaskel-Rs, we analyzed by flow cytometry the expression level of Gaskel-Rs that are capable of binding Gaskel, on the surface of resting platelets from WT mice or mice lacking any 1 of the Gaskel-Rs. Platelets were incubated with recombinant myc-tagged Gaskel (Gaskel-myc), and binding of Gaskel-myc was measured by flow cytometry using an anti-myc antibody. As shown in Figure 10, Gaskel-myc was found to bind to WT platelets, while only negligible amounts of Gaskel-myc bound to Tyro3–/–, Axl–/–, or Mer–/– platelets. These findings thus reveal that, in the absence of 1 Gaskel-R subtype, the expression of the other Gaskel-Rs is also minimal. To confirm that the expression of the Gaskel-Rs was indeed regulated in an interdependent manner, we measured by flow cytometry, using an anti-Axl antibody, the expression level of Axl (as a prototype of the Gaskel-Rs) on platelets.

ATP) in response to 2 μg/ml collagen was significantly impaired in Tyro3–/– platelets (9.9 ± 0.7 μM ATP from WT platelets versus 5.3 ± 0.2 μM ATP from Tyro3–/– platelets; n = 3, P < 0.05). Secretion of α granules, assessed by measurement of the surface levels of P-selectin during platelet activation, was also impaired in Tyro3–/–, Axl–/–, or Mer–/– platelets in response to low concentrations of ADP (Figure 8). Thus, defects in granule secretion may also contribute to the phenotypes of Gaskel-R deficiency.

Figure 5
Effect of the lack of Tyro3, Axl, or Mer on fibrinogen binding. Loss of 1 Gaskel-R (results shown for Axl–/– mice) does not inhibit fibrinogen binding to αIIbβ3 integrin. The binding of labeled fibrinogen to platelets activated with ADP (1.25–20 μM) was measured by flow cytometry. Black lines denote resting platelets; red shading denotes ADP-stimulated platelets. A representative example of 4 independent experiments is shown.
As expression of Axl on the surface of resting platelets was below the detection limit of this assay (data not shown), we stimulated platelets with the PAR4-activating peptide (0.25 mM). Activated WT platelets expressed detectable levels of Axl on their surface, while, as expected, Axl was absent from activated Axl−/− platelets (Figure 11). Notably, however, activated Tyro3−/− and Mer−/− platelets expressed only minimal amounts of Axl on their surface (Figure 11). Similar results were obtained from analysis of the expression of Tyro3 (platelets expressing Tyro3 in WT mice, 77% ± 4%; in Tyro3−/− mice, 6% ± 0.5%; in Axl−/− mice, 38% ± 6%; and in Mer−/− mice, 34% ± 4%; mean ± SEM, n = 3, P < 0.05) or Mer (platelets expressing Mer in WT mice, 70% ± 3%; in Tyro3−/− mice, 39% ± 2%; in Axl−/− mice, 35% ± 3%; and in Mer−/− mice, 9% ± 0.6%; mean ± SEM, n = 4, P < 0.05). Thus, lack of any 1 of the Gas6-Rs significantly reduced the expression of the other Gas6-Rs on the surface of platelets and, thus, the capacity to bind and respond to activation by Gas6.

Gas6 signaling activates PI3K and Akt. The signaling pathway, downstream of Gas6-R activation, was further investigated. The Gas6 pathway does not seem to influence the eicosanoid synthesis pathway, as Gas6−/− platelets produce normal amounts of TXA2 (7) and Gas6-R signaling in other cell types requires PI3K and Akt (18–20). We therefore analyzed whether Gas6-R signaling might activate PI3K and Akt, as the PI3K pathway is involved in αIIbβ3 activity regulation in platelets (4, 29). Indeed, Gas6 was found to enhance phosphorylation of PI3K and Akt in resting WT platelets, but not in Tyro3−/− (Figure 9A), Axl−/−, or Mer−/− (data not shown) platelets, indicating that each Gas6-R is critical for signal transduction of Gas6. The Akt pathway is also well known to play a key role in response to thrombin. In WT platelets, Akt phosphorylation was already increased within 30 seconds after activation with 0.05 IU/ml thrombin (Figure 12). In contrast, Akt phosphorylation in Tyro3−/−, Axl−/−, (data not shown), and Mer−/− (Figure 12, A and B) platelets occurred only after 3 minutes, and only when the platelets were stimulated with a 10-fold higher concentration of thrombin (0.5 IU/ml; Figure 12, A and B). Thus, phosphorylation of Akt was impaired in platelets lacking any 1 of the Gas6-Rs.

**The soluble Axl extracellular domain inhibits platelet function.** We next assessed the therapeutic potential of blocking the Gas6-R-dependent amplification of outside-in signaling via αIIbβ3 by using the soluble Axl extracellular domain (hAxl-EC-Fc), which traps Gas6. The aggregation of WT platelets in response to ADP (5 μM) was dose-dependently blocked by hAxl-EC-Fc. In contrast, hAxl-EC-Fc had no effect on WT platelets in the absence of ADP (Figure 3D). Administration of 0.75 mg/kg hAxl-EC-Fc significantly protected WT mice against fatal collagen/epinephrine-induced pulmonary thromboembolism. Indeed, 60% of the mice receiving hAxl-EC-Fc survived, whereas none of the mice receiving control vehicle survived (n = 7, P < 0.05). Mice treated with hAxl-EC-Fc did not spontaneously bleed, nor were their bleeding times prolonged (data not shown). These results indicate that inhibition of Gas6-R signaling by trapping of its ligand effectively prevents thrombosis.
cal thrombosis in vivo; (b) Gas6-R signaling appears to affect outside-in signaling via the $\alpha_{\text{III}}\beta_3$ integrin and to regulate granule secretion; and (c) activation of Gas6-R by Gas6 activates PI3K and stimulates tyrosine phosphorylation of $\beta_3$ integrin. These genetic studies provide novel insight into the poorly understood physiological importance of outside-in signaling, which is critical for the formation of stable platelet plugs and clot retraction.

Gas6-R$^{-/-}$ mice are phenocopies of Gas6$^{-/-}$ mice. Mice lacking any 1 of the 3 Gas6-Rs (Tyro3, Axl, or Mer) have normal initial bleeding times and are resistant to thrombosis, but they do not suffer spontaneous bleeding. Their resistance to thrombosis was not due to impaired megakaryocytopenesis or defects in the coagulation cascade but was due, at least in part, to platelet dysfunction. As Gas6 seems to regulate also the expression of tissue factor on activated endothelial cells (E.M. Conway and P. Carmeliet, unpublished observations), we cannot exclude the possibility that Gas6-Rs also contribute to thrombus formation via such a mechanism. However, the platelet transfusion experiments clearly indicate that the antithrombotic phenotype in Gas6-R–deficient mice is, at least in part, attributable to a platelet defect. In the absence of a Gas6-R, low concentrations of ADP, collagen, or TXA2 induced reversible platelet aggregation, while high concentrations of these platelet agonists caused irreversible platelet aggregation. In addition, low concentrations of thrombin or the PAR4-activating peptide stimulated platelet aggregation. Thus, aggregation of platelets lacking any 1 of the Gas6-Rs is comparable to that of aspirin, when assayed in the same thrombembolism model (30).

A remarkable finding was, however, that deficiency of any of the Gas6-Rs protected mice against thrombosis to the same significant extent, which suggests that each of these receptors has a comparably important role in platelet aggregation. Even more, Gas6 was capable of inducing tyrosine phosphorylation of each Gas6-R, but absence of 1 Gas6-R prevented tyrosine phosphorylation of the 2 other Gas6-Rs. An explanation for the latter phenomenon is provided by the observation that substantial amounts of Gas6-R reached the platelet surface only during platelet activation and that the lack of any 1 of the Gas6-Rs dramatically reduced the surface expression of the other receptors. As a result, the binding of Gas6 to Gas6-Rs on the platelet surface was significantly reduced to negligible residual levels when 1 of these Gas6-Rs was absent. These findings suggest that Gas6-Rs somehow cross-talk to each other to codetermine their surface exposure and responsiveness to Gas6 (i.e., binding of ligand, stimulation of receptor tyrosine kinase activity). Thus, the exposure of any 1 of the Gas6-Rs appears to

**Figure 8**

Effect of the absence of Tyro3, Axl, or Mer on P-selectin expression, a marker of granule secretion. Analysis by flow cytometry revealed that P-selectin levels on the platelet surface were significantly reduced in Tyro3$^{-/-}$, Axl$^{-/-}$, or Mer$^{-/-}$ platelets, after activation with ADP. Black lines denote resting platelets; red shading denotes ADP-stimulated platelets. A representative example of 3 independent experiments is shown.

**Figure 9**

Gas6 signaling through its receptors Tyro3, Axl, and Mer. (A) Gas6 promotes phosphorylation of its receptors (shown for Axl), PI3K, Akt, and $\beta_3$ integrin in WT but not in Tyro3$^{-/-}$, Axl$^{-/-}$ (not shown), or Mer$^{-/-}$ (not shown) platelets. Platelets were incubated with 400 ng/ml human recombinant Gas6 (hrGas6) for 3 minutes. For detection of Axl or PI3K phosphorylation, platelets were lysed and phosphotyrosine-containing proteins were immunoprecipitated. The precipitates were analyzed by SDS–PAGE and Western-blotted with anti–phospho-Akt antibody, anti–total Akt antibody, or anti–phospho-PI3K antibody. B, WT and Tyro3$^{-/-}$ platelets were stimulated with increasing concentrations of thrombin. Black lines denote controls; red shading denotes platelets stained with PE-conjugated anti–$\beta_3$ integrin antibody. C, WT and Tyro3$^{-/-}$ platelets were stimulated with increasing concentrations of thrombin for 3 minutes. A representative example of 3 independent experiments is shown.
Deficient platelets were able to overcome their defect when stimulated to rebleed after initial cessation of bleeding. A difference between clot retraction upon thrombin treatment and showed a tendency of partial aggregation in response to ADP. Both strains also had defective binding Tyro3, Axl, or Mer bound normal amounts of fibrinogen in response to ADP. While these findings are consistent with a model whereby Gas6, secreted by activated platelets, amplifies fibrinogen-induced platelet spreading, irreversible platelet aggregation, and clot retraction by inducing signals that lead to phosphorylation of Gas6-Rs, which in turn activate PI3K and Akt. Flow serine and threonine phosphorylation, and subsequently tyrosine phosphorylation of the cytoplasmic tail of β3 integrin, are precisely modulated via Gas6-R activation remains to be further investigated but might involve intermediate activation of PI3K and Akt in association with integrin-linked kinase displacement toward αIIbβ3 (4).

Absence of a Gas6-R impairs outside-in signaling. Platelets lacking Tyro3, Axl, or Mer bound normal amounts of fibrinogen in response to ADP. While these findings are consistent with a model whereby Gas6-R signaling would not be necessary for initial activation of αIIbβ3 by inside-out signaling, the present data do not exclude the possibility that Gas6-R signaling might modulate this process at a later stage. In fact, our data indicate that the Gas6-R signaling was required for tyrosine phosphorylation of the β3 integrin and thus for outside-in αIIbβ3 signaling. Such a model also explains why mice lacking any 1 of the Gas6-Rs had a tendency to rebleed after transient hemostasis and had delayed platelet spreading and clot retraction, processes consistent with a defect in regulation of outside-in αIIbβ3 signaling (1). The platelet dysfunction caused by absence of a Gas6-R resembles that of mice in which the cytoplasmic tyrosine residues of the β3 integrin subunit were mutated to phenylalanine (DiYF mice) (2). Both DiYF mice and Gas6-R-deficient mice exhibit normal fibrinogen binding and a normal first wave of ADP-induced platelet aggregation. In contrast, outside-in signaling in DiYF and Gas6-R-deficient platelets was impaired, as they spontaneously disaggregated following initial aggregation in response to ADP. Both strains also had defective clot retraction upon thrombin treatment and showed a tendency to rebleed after initial cessation of bleeding. A difference between the DiYF and the Gas6-R-deficient phenotypes is that Gas6-R-deficient platelets were able to overcome their defect when stimulated by a high dose of platelet agonists for a long duration; this indicates that the absence of a Gas6-R caused a milder outside-in signaling defect than the DiYF mutation did. Our findings obviously do not exclude the possibility that Gas6-R signaling might be involved in additional mechanisms, regulating platelet plug stability and clot retraction. In fact, our data that Gas6-Rs also regulate the process of secretion from dense and α platelet granules and activate PI3K and Akt could imply that perpetuation of the platelet plug might be amplified by upregulation of β3 integrin activity, leading to enhanced platelet aggregate stability.

Gas6-R signaling. Activation of the Gas6-Rs stimulated phosphorylation of PI3K and Akt (34). PI3K plays an important role in strengthening platelet aggregation during the irreversible phase of aggregation (35, 36). Taken together, our data are consistent with a model whereby Gas6, secreted by activated platelets, amplifies fibrinogen-induced platelet spreading, irreversible platelet aggregation, and clot retraction by inducing signals that lead to phosphorylation of Gas6-Rs, which in turn activate PI3K and Akt. Flow serine and threonine phosphorylation, and subsequently tyrosine phosphorylation of the cytoplasmic tail of β3 integrin, are precisely modulated via Gas6-R activation remains to be further investigated but might involve intermediate activation of PI3K and Akt in association with integrin-linked kinase displacement toward αIIbβ3 (4).

Gas6/Gas6-R signaling: a novel class of platelet modifiers. Deficiency of the β3 integrin (37, 38), of either of the purinergic receptors P2Y1 (39, 40) and P2Y12 (41), of the GTP-binding G protein (42), or of protease-activated receptor 3 (PAR3) or PAR4 (26, 43) protects mice against thrombosis but also causes bleeding. Deficiency of Gas6 or any of its receptors also protected mice against thrombosis but did not prolong the bleeding time after tail clamping. In this respect, the effect of Gas6-R signaling resembles that of Eph kinases and ephrins (5) and CD40 ligand (44). Whereas Gas6 by itself is not able to promote platelet aggregation (7), Gas6 signal-
ing is critical for platelet function during the late phase of platelet activation, when platelets come into close contact with each other. Gas6 perpetuates platelet activation by enhancing PI3K, Akt, granule secretion, and β3 integrin signaling. Gas6 thus belongs to a class of molecules that are redundant for base-line hemostasis but that constitute an important “amplification” system in pathological conditions. Abnormal Gas6 signaling through 1 or several of its receptors might contribute to bleeding disorders due to a platelet dysfunction. Conversely, abnormal activation of the Gas6 pathway might also contribute to disease phenotypes due to platelet activation (e.g., heparin-induced thrombocytopения, anti-phospholipid antibody syndrome) that are characterized by thrombosis. A very recent study indicates a significant association between a Gas6 polymorphism and ischemic stroke, placing Gas6 as a promising candidate for a susceptibility gene for this disease (45).

Medical implications. Based on the critical role of c-terminal β3 in platelet function, pharmacological inhibitors of ligand binding to c-terminal β3 have been developed to prevent thrombosis. With the exception of recombinant antibodies that block fibrinogen binding to c-terminal β3, and therefore aggregation as well as outside-in signaling, all currently available antiplatelet drugs are specific inhibitors of inside-out signaling. For example, aspirin inhibits COX-1 and, consequently, the production of TXA2, while clopidogrel antagonizes signaling through the P2Y12 receptor. Drugs designed to prevent irreversible platelet aggregation rather than inhibition of formation of platelet aggregates are currently not available. Our findings indicate that inhibition of Gas6, involved in the signaling pathways controlling irreversible aggregation, could be a novel treatment paradigm. Indeed, a soluble Axl extracellular domain protected mice against fatal thromboembolism without bleeding side effects. Axl extracellular domain therefore provides a novel and relatively safe means to prevent thrombosis.

Methods

Materials. Busulfan, ADP, U46619, and A23187 were from Sigma-Aldrich. Collagen was from Hormon Chemie. The PAR4 agonist peptide AYPGKF was synthesized by Bachem AG.

Mice. Tyro3−/−, Axl−/− and Mer−/− mice were progeny of the original colony of 50% 129/50% C57BL/6 mice (21). WT mice used as controls had the same genetic background. Ethical approval for the animal experiments detailed in this article was received from the Institutional Animal Care and Research Advisory Committee of the University of Leuven and the Ethical Committee for Animal Experiments of the Medical Faculty of the University of Geneva.

Hemostasis. We measured bleeding by 2-mm tail-tip transection in mice anesthetized by i.p. injection of a mixture of ketamine (80 mg/kg) and xylazine (16 mg/kg). We used 2 models of bleeding time. For the first model, a stopwatch was started immediately upon transection to determine time to cessation of bleeding. Blood drops were removed every 30 seconds using a filter paper. If bleeding did not recur within 60 seconds of cessation, bleeding was considered stopped. The second model was used to detect rebleeding after initial hemostasis. For quantitative estimation of the amount of blood loss per time unit after tail transection, the cut end of the tail was immediately immersed in 20 ml normal saline at 37°C. Aliquots of saline solution containing blood were sampled every minute for 10 minutes and then every 5 minutes until 30 minutes. Red blood cells were lysed using Zap-Oglobin II lytic reagent (Beckman Coulter Inc.), and samples were read at 405 nm (to measure the hemoglobin content). A quantitative estimation of the amount of blood loss was determined by measurement of the hemoglobin content of aliquots of blood collected in saline after tail transection as compared with known volumes of blood from the same mouse diluted in the same amount of saline.

Human recombinant Gas6 and Axl extracellular domain. Human recombinant Gas6 was produced in a human embryonic kidney cell line (HEK293) stably transfected with the pSecTag2/Hygro expression vector (Invitrogen Corp.) encoding the human Gas6 cDNA. Human recombinant Gas6 was purified from conditioned medium using a modification of a calcium affinity method used for proteins containing γ-carboxy-glutamic acid residues (47, 48). The cDNA encoding a 415-AA mature human Axl extracellular domain (hAxl-EC) was cloned from a human kidney cDNA library (Stratagene). Splice overlap extension was used to fuse the cDNA downstream of the secretion signal of the pSecTag2/Hygro vector (Invitrogen Corp.), followed by in-frame sequence coding for the human IgG-Fc domain (R&D Systems Inc.). The resulting expression vector drives expression and secretion of a hAxl-EC-Fc fusion under the control of the CMV promoter. The plasmid pSecTag2/Hygro–hAxl-EC-Fc was stably transfected in HEK293 cells. The hAxl-EC-Fc was purified on protein A-Sepharose.

Thrombus models. Mice were anesthetized by i.p. injection of a mixture of ketamine (80 mg/kg) and xylazine (16 mg/kg). Thrombus formation due to stasis was induced by injection of tissue thromboplastin (1 vol Innovin/49 vol saline; Innovin was from Dade Behring Inc.) into the dorsal vein of the penis (5 ml/kg), followed immediately by tightening of I suture in the inferior vena cava just below the left renal vein for 15 minutes. Thrombosis was quantified by weighing of the thrombus (49). In mice anesthetized by i.p. injection of sodium pentobarbital (60 mg/kg), thromboembolism was induced by injection of a mixture of collagen (0.5 mg/kg, equine collagen; Hormon Chemie) and epinephrine (60 μg/kg) into the jugular vein (30). When indicated, mice received 10 μg hAxl-EC-Fc.

Platelet aggregation. Platelet aggregation was performed as described in ref. 7. Briefly, whole blood, drawn from anesthetized mice from the inferior vena cava into 3.13% citrate (1 vol anticoagulant/9 vol blood), was centrifuged at 100 g (10 minutes) to obtain PRP and additionally at 2,000 g (10 minutes) to obtain platelet-poor plasma (PPP). PRP and PPP were pooled from 4–6 mice. To prepare washed platelets, asprarye was added to PRP (final concentration, 1 IU/ml), and platelets were incubated for 10 minutes at 37°C, washed by addition of 2 vol acid-citrate-dextrose solution (22 g trisodium citrate dihydrate, 8 g citric acid monohydrate, and 25 g glucose dextrose in 1 liter of H2O2), and centrifuged at 2,000 g (10 minutes). The platelet pellet was resuspended in Tyrode’s buffer (pH 7.3) containing 137 mM NaCl, 2.7 mM KCl, 12 mM NaHCO3, 0.4 mM NaH2PO4, 5 mM HEPES, and 0.1% BSA. CaCl2 (2 mM final) and MgCl2 (1 mM final) were added immediately before platelet stimulation. Platelet aggregation was analyzed using a Multiplate (Dade Behring).
measured turbidimetrically using an optical Platelet Aggregation Profiler (model PAP-4; Bio/Data Corp.). When indicated, WT washed platelets were preincubated with hAxl-EC-Fc before stimulation with ADP (1 minute). Clot retraction. PRP was diluted with PPP to bring the platelet count to 100 G/l and put into an aggregation cuvette. Thrombin was added at a final concentration of 10 IU/ml. The aggregation cuvette was then incubated at 37°C in the aggregeometer for 2 hours without stirring, while the clot retraction was monitored every 15 minutes (N. Prévost, unpublished observations).

Fibrinogen binding. Washed platelets (10^6/ml in final volume) were incubated simultaneously with a mixture of PBS (negative control) or ADP (1.25-20 μM in final volume), Oregon Green 488-conjugated fibrinogen (150 μg/ml in final volume; Invitrogen Corp.), and Tyrode’s buffer containing 0.1% BSA in a final volume of 200 μl. After 10 minutes at 37°C under stirring conditions, samples were fixed by the addition of paraformaldehyde (1% in final volume). Platelets were then resuspended in 500 μl PBS containing 0.2% BSA, and samples were analyzed by flow cytometry using a FACSscan instrument (BD Biosciences) and CellQuest software (BD Biosciences).

Platelet secretion. Dense-granule secretion, investigated by platelet ATP release in response to collagen, was monitored in PRP by addition of firefly luciferase and luciferin and comparison of the luminescence generated by platelet ATP release with that generated by an ATP standard (CHRONOLUME; Chrono-Log Corp.), as reported previously (7). To measure α-granule secretion, washed platelets (10^6/ml) were incubated with ADP at various concentrations for 10 minutes at 37°C under stirring conditions and subsequently labeled with FITC-conjugated anti–P-selectin antibody (BD Biosciences—Pharmingen) and analyzed by flow cytometry using a FACSscan instrument and CellQuest software, as described previously (7).

Adhesion to immobilized fibrinogen. Adhesion to immobilized fibrinogen was quantified as described previously (50). Human fibrinogen (Sigma-Aldrich) in PBS (pH 8.0) was plated onto microtiter plates at concentrations ranging from 1 ng to 2 μg per well and incubated overnight at 4°C. Plates were washed twice with PBS (pH 7.4) and blocked for 2 hours at room temperature with 20 μl PBS containing 0.1% BSA. After 1 hour of incubation, nonadherent platelets were removed, and the wells were washed twice with 150 μl Tyrode’s buffer supplemented with 1 mM MgCl2. One hundred fifty microliters of pNpp buffer (0.1 M citrate [pH 5.4], 0.1% Triton X-100, 5 mM p-nitrophenylphosphate) was added for 1 hour at room temperature. Then, 100 μl of 2-M NaOH was added, and adherent platelets were quantified in a microplate reader at 405 nm. The percentage of platelets adhering was determined by calculation of the ratio of bound to maximal signal at 405 nm, where maximal reading was obtained from a microtiter well containing 6 × 10^6 platelets that was not subjected to washing procedures.

Rhodamine-phalloidin staining. A Lab-Tek chamber slide (Nalge Nunc International) was coated with 100 μg/ml fibrinogen. Washed platelets (6 × 10^6) in Tyrode’s buffer supplemented with CaCl2 and MgCl2 were added to each well of the chamber slide for 15–60 minutes at 37°C. Nonadherent platelets were washed away, and adherent cells were fixed in 1% paraformaldehyde in PBS for 15 minutes at room temperature. After 3 washes with PBS, excess aldehyde was quenched with 10 mM ethanolamine in PBS (or 0.1 M glycine in PBS) for 5 minutes. Platelets were permeabilized in 0.1% Triton X-100 in PBS for 1 minute and incubated in rhodamine-phalloidin (Sigma-Aldrich) diluted 1:100 in PBS for 15 minutes. The slide was removed from the chamber and then washed 3 times in PBS. Slides were then mounted in VECTASHIELD (Vector Laboratories Inc.). Platelet spreading was quantified using an Axioptophor microscope equipped with an AxioCam camera (Carl Zeiss AG).

Platelet spreading by video microscopy. Observation chambers coated with human fibrinogen in PBS (pH 8.0) (2 mg/ml) were washed twice with PBS (pH 7.4) and blocked for 2 hours at room temperature with 20 mg/ml BSA in PBS. Then, washed platelets resuspended in Tyrode’s buffer (10^6) were dropped into the chambers and observed with an Axiovert 100 TV inverted microscope (Carl Zeiss AG), equipped with a heated stage (37°C) and a digital charged-coupled device camera (C4742-95-10; Hamamatsu Photonics). Platelet spreading was recorded with Openlab 3.0.6 software (Improvision) with frame intervals of 15–30 seconds.

Immunoprecipitation and Western blotting. For immunoprecipitation, washed platelets were lysed in the presence of the following protease inhibitors: 1 μM NaVO3, 1 mM NaF, and complete inhibitors (Roche Diagnostics Corp.). Insoluble materials were removed by centrifugation for 10 minutes at 4°C. The resulting supernatants were immunoprecipitated with monoclonal anti-phosphotyrosine–agarose (clone PT-66) beads (Sigma-Aldrich). For dose-response and time-course experiments, stirred washed platelets (4 × 10^6 platelets in 100 μl Tyrode’s buffer) were stimulated with human recombinant Gas6 or thrombin at 37°C. The reaction was stopped by the addition of SDS sample buffer. Sample aliquots were loaded on SDS-PAGE (10%) and subjected to Western blotting using 1 of the following antibodies: anti-Rse (also called Tyro3), anti-Axl, anti-Mer, anti–PI3K p85α, anti–β3 integrin (Santa Cruz Biotechnology Inc.), the PhosphoPlus Akt (Ser473) antibody (Cell Signaling Technology Inc.), or anti–β3 integrin [pY773] phospho-specific antibody (BioSource International Inc.). The data are representative of 3–4 independent experiments using PRP pooled from 6 WT, Tyro3–/–, Axl–/–, or Mer–/– mice.

Detection of β3 integrin on platelet surface by flow cytometry. Washed platelets (10^6/ml) were incubated with PE-conjugated anti–β3 integrin antibody (BD Biosciences—Pharmingen) and analyzed by flow cytometry using a FACSscan instrument and CellQuest software.

Gas6 binding. Washed platelets (10^6/ml in final volume) were incubated simultaneously with PBS (negative control) or myc-tagged recombinant Gas6 (5 μg/ml in final volume) and FITC-conjugated anti–myc-tag antibody (Abcam Ltd. in Tyrode’s buffer containing 0.1% BSA in a final volume of 200 μl at 37°C for 10 minutes under stirring conditions. The reaction was stopped by the addition of paraformaldehyde (1% in final volume), and platelets were resuspended in 500 μl PBS containing 0.2% BSA and analyzed by flow cytometry using a FACSscan instrument and CellQuest software.

Detection of Gas6-R on platelet surface by flow cytometry. Resting washed platelets (10^6/ml in final volume) or washed platelets activated by the PAR4-activating peptide AYPGKF at 0.25 mM for 10 minutes at 37°C under stirring conditions were incubated with an anti-Tyro3, an anti-Axl, or an anti-Mer antibody (R&D Systems Inc.). Tyro3, Axl, or Mer was subsequently detected on platelets by a FITC-conjugated rabbit anti-rat antibody and analyzed by flow cytometry using a FACSscan instrument and CellQuest software.

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Address correspondence to: Anne Angelillo-Scherrer, Division of Angiology and Hemostasis, University Medical Center, 1 rue Michel-Servet, CH-1211 Geneva 14, Switzerland. Phone: 41-22-379-55-67; Fax: 41-22-372-92-99; E-mail: Anne.Angelillo@medecine.unige.ch.


