Central role of the P2Y\textsubscript{12} receptor in platelet activation

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Platelet activation occurs in response to vessel injury and is important for the arrest of bleeding. Platelet activation during disease states leads to vascular occlusion and ischemic damage. The P2Y\textsubscript{12} receptor, activated by ADP, plays a central role in platelet activation and is the target of P2Y\textsubscript{12} receptor antagonists that have proven therapeutic value.


The vessel wall contains a continuous lining of endothelium that serves as a barrier between the circulating platelets and the prothrombotic subendothelial matrix (1). Upon vessel injury, the endothelial layer is disrupted and the circulating platelets are exposed to subendothelial proteins such as vWF, collagen, and vitronectin, among others (1). The platelets initially interact with the subendothelium through adhesive receptors, such as GP\textsubscript{Ib-IX-V} receptors, that mediate rolling and tethering of the platelets to vWF at the site of vascular injury.

Next, the platelet collagen receptors \(\alpha_2\beta_1\) and GP\textsubscript{VI} mediate a more firm adhesion and cause further platelet activation. These initial interactions with the subendothelium cause the release of contents from the platelet dense granules, which contain platelet agonists such as ADP, and the \(\alpha\)-granules, which contain fibrinogen, factor V, and P-selectin (1). The release of the granule contents causes further platelet activation, but it also fuels the coagulation response as a result of the release of factor V and fuels the inflammatory response through the exposure of P-selectin on the platelet surface. The platelet also generates lipid mediators such as thromboxane A\textsubscript{2}, ADP elicits its effects on the platelet through the P2Y\textsubscript{1} and P2Y\textsubscript{12} receptors (2), whereas thromboxane A\textsubscript{2} activates the thromboxane-prostanoid (TP) receptor on the platelet surface (1). The released dense granule contents cause further platelet activation and recruitment of circulating platelets to the site of injury. Platelets interacting with these mediators also undergo platelet shape change, a process of actin cytoskeletal reorganization that changes the platelets from a disc shape to a round shape with long, filopodial extensions that form a meshwork of platelets in the platelet plug (3). Also, tissue factor is exposed, which initiates the coagulation response that results in formation of thrombin. Thrombin activates platelets via interactions with the proteinase-activated receptor-1 (PAR1) and PAR4 receptors (4) and also cleaves fibrinogen to form fibrin. Fibrin further stabilizes the accumulating platelet plug at the site of injury, resulting in a stable hemostatic plug.

Interactions of the platelets with collagen, vWF, ADP, thromboxane A\textsubscript{2}, and thrombin cause intracellular platelet signaling that leads to the activation of the heterodimeric integrin \(\alpha_{IIb}\beta_3\), also known as the fibrinogen receptor (5). The intracellular platelet signaling from these agonists causes the fibrinogen receptor to change from a low-affinity state to a high-affinity state that binds fibrinogen (6). Fibrinogen binds to the platelets via the activated fibrinogen receptor, and this cross-linking of platelets to fibrinogen results in platelet aggregates that accumulate and arrest bleeding at the site of injury (Figure 1). Thus, platelet activation is the product of many signals originating from many receptors, which each contribute to the formation of a platelet plug.

Pathophysiologic conditions, such as atherosclerotic plaque rupture, can lead to aberrant platelet activation resulting in arterial thrombosis, which can cause myocardial infarction and ischemic stroke (6). The importance of ADP in this process has been demonstrated both by antiplatelet drugs that target the P2Y\textsubscript{12} receptor (2) and by patients with dysfunctional P2Y\textsubscript{12} receptors (7). Antagonism of the P2Y\textsubscript{12} receptor with either ticlopidine or clopidogrel is clinically effective in the prevention of myocardial infarction, ischemic stroke, and vascular death (8). Despite the established role of the P2Y\textsubscript{12} receptor in the hemostatic response,
the full implications of P2Y12 receptor antagonism in the prevention of thrombosis remain incompletely understood. It is hoped that more clinically effective P2Y12 antagonists will prevent the incidence of ischemic events that stem from aberrant platelet activation and therefore will be used as improved and suitable treatments for thrombosis.

The central role of the P2Y12 receptor: ex vivo effects

Prior to the cloning of the P2Y12 receptor, drugs that selectively target this receptor had been widely used as antplatelet agents (2). Ex vivo studies used platelets treated with clopidogrel or reversible antagonists of the P2Y12 receptor and led to the conclusion that the P2Y12 receptor is crucial to several platelet functions. Thus far, studies have identified a potentiating role for the P2Y12 receptor in dense granule secretion (9), fibrinogen-receptor activation (10–14), and, as reported in a recent issue of the JCI, thrombus formation (15, 16), identifying it as a central mediator of the hemostatic response. This receptor is also important for the irreversible platelet aggregation induced not only by ADP, but also by thromboxane A2 and the PAR1-selective peptide agonist SFLLRN (17, 18). The P2Y12 receptor also causes inhibition of stimulated adenylyl cyclase (19, 20) but does not play any role in ADP-induced platelet shape change and intracellular calcium mobilization (17, 18). Furthermore, G1 signaling that is mediated by P2Y12 receptor activation can lead to platelet aggregation when either Gq or G12/13 pathways are simultaneously stimulated (10, 11, 14), or by itself when exposed to high concentrations of ADP (100 µM) (21). The P2Y12 receptor plays a crucial role in ADP-mediated generation of thromboxane A2, another important platelet activator (22). Signaling events downstream of the P2Y12 receptor also potentiate agonist-induced dense granule release and procoagulant activity (17). In addition, α-granule release and subsequent expression of P-selectin on activated platelets

Figure 1

The hemostatic process. Upon vessel injury, platelets roll and become tethered to the vessel wall by interactions with vWF and collagen (noted as black strands). These interactions cause platelet shape change, and release of ADP from dense granules. The activated platelet also generates thromboxane A2 (TxA2). Both ADP and TxA2 are agonists that cause further platelet activation and accumulation of platelets at the site of injury. Vessel injury also causes exposure of tissue factor, which catalyzes the coagulation response. This response results in the formation of thrombin, which further activates platelets and cleaves fibrinogen to form fibrin. The combination of activated platelets and fibrin at the site of injury forms a stable hemostatic plug that arrests bleeding.
depend on P2Y12 activation (23, 24). Interestingly, all the functions of the P2Y12 receptor can be mimicked by epinephrine, which stimulates members of the G\(_i\) family of G proteins by binding to the platelet \(\alpha_2\) receptor (17). Thus, the P2Y12 receptor plays a central role in platelet activation, in the recruitment of other platelets to the site of injury subsequent to the adhesion of platelets to vWF and collagen, and in the enhancement of the efficiency of platelet activation by other agonists such as thrombin and thromboxane A\(_2\), which are generated as secondary platelet agonists (Figure 2).

The P2Y12 receptor couples primarily to G\(_{\alpha_i2}\) and less prominently to other members of the G\(_i\) family, resulting in the inhibition of adenylyl cyclase (25). Epinephrine, through stimulation of the \(\alpha_2\) receptor and resulting G\(_i\) signaling, also achieves the same effect (26). However, reduced levels of cAMP are not directly responsible for the downstream effects of P2Y12 receptor activation (25, 26). Gi signaling leads to activation of PI3K, Akt, Rap1b, and potassium channels (17). Mice lacking PI3K-\(\gamma\) show aberrations in platelet function only when low doses of ADP are used, but are provided protection from thromboembolism (27). Recent studies indicate that Rap1b, Akt, and potassium channels are important functional effectors downstream of P2Y12 receptor stimulation (28–31) (Figure 3).

**Patients with defective P2Y12 receptor function**
Patients with a defect in the gene encoding the P2Y12 receptor have a congenital bleeding disorder (7, 32–35). A number of patients have been identified that have decreased aggregation responses to ADP and to low doses of other agonists such as collagen and thrombin (32–35). These patients generally have normal platelet shape change responses to ADP but have impaired abilities to inhibit adenylyl cyclase activity (33). While patients with defective P2Y12 receptor function have dense granules that are normal in both numbers and content, platelet release of granules is generally decreased because of the potentiating effects of the P2Y12 receptor on granule secretion (34). Patients and mice lacking functional P2Y12 receptors have increased bleeding times (7, 32, 36). Patients who are heterozygous for the P2Y12 receptor bind intermediate amounts of 2-methylthio-ADP (2-MeSADP), an agonist of the P2Y1 and P2Y12 receptors, and also have extended bleeding times (7, 34, 37). Some patients with P2Y12 receptor deficiency have been shown to possess a truncated form of the receptor due to deletions in the gene, whereas other individuals have mutations that lead to impaired P2Y12 receptor function (38). Analysis of the P2Y12 receptor sequence from a patient with impaired ADP responses has identified amino acid residues important in P2Y12 receptor function (35). A G-to-A mutation in one allele changed Arg256 into Gln, while a C-to-T alteration resulted in Arg265 changing to Trp (35). Though receptor number
Table 1
In vivo effects of P2Y12 receptor blockade

<table>
<thead>
<tr>
<th>Species</th>
<th>P2Y12 blockade</th>
<th>Injury/stimulus</th>
<th>Embolization</th>
<th>Bleeding time</th>
<th>In vivo thrombus characteristics</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rabbit (mesenteric arterioles)</td>
<td>AR-C69931MX</td>
<td>Mechanical injury</td>
<td>↓</td>
<td>Microvessel bleeding time unchanged</td>
<td>Size of thrombus reduced</td>
<td>15</td>
</tr>
<tr>
<td>Mouse (mesenteric artery)</td>
<td>P2Y12+/−</td>
<td>Ferric chloride, tail bleeding</td>
<td>↑</td>
<td>Tail bleeding time extended, ↑ time to occlusion</td>
<td>Delayed, unstable, nonocclusive thrombus</td>
<td>16</td>
</tr>
<tr>
<td>Mouse</td>
<td>P2Y12+/−</td>
<td>Ferric chloride, tail bleeding</td>
<td>Unchanged from wild type</td>
<td>Unchanged from wild type</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Human Homozygous</td>
<td></td>
<td></td>
<td></td>
<td>Extended from normal</td>
<td>ND</td>
<td>32-35</td>
</tr>
<tr>
<td>Human Heterozygous</td>
<td></td>
<td></td>
<td></td>
<td>Extended from normal</td>
<td>ND</td>
<td>34</td>
</tr>
</tbody>
</table>

The role of the P2Y12 receptor in thrombus formation and its effect on bleeding times have been characterized using pharmacologic antagonists, mice deficient in the P2Y12 receptor, and patients who have a mutation or truncation of the P2Y12 receptor. Multiple species and models of thrombosis have been used to characterize the effect of P2Y12 blockade in vivo, leading to varying conclusions and interpretations. ND, not determined.

and affinity for 2-MeSADP were unchanged, these mutations were demonstrated to lead to impaired Gi signaling in response to P2Y12 stimulation (35). Expressed P2Y12 receptors containing these mutations had a similar loss of function (35). Thus, these residues play an important role in the function of the P2Y12 receptor but have no effect on the binding of ADP.

Bleeding times
The P2Y12 receptor antagonist clopidogrel has been shown to be efficacious when occupying even 50% of the P2Y12 receptor population (39); thus an antagonist of the P2Y12 receptor could be an effective therapeutic even when 50% of the P2Y12 receptors are functioning. Patients deficient in the P2Y12 receptor, such that their platelets bind only intermediate levels of 2-MeSADP, have slightly prolonged bleeding times (7, 34, 37). The P2Y12 receptor has been cloned, and mice that are deficient in the P2Y12 receptor have been generated. Consistent with observations in patients deficient in the P2Y12 receptor, mice lacking the P2Y12 receptor have increased tail bleeding times (19, 36, 40, 41). However, heterozygous mice show little or no change in bleeding times (16). It is currently unclear why a 50% decrease in the population of functional P2Y12 receptors, either following clopidogrel treatment or in humans heterozygous for the P2Y12 receptor, prolongs bleeding time, while heterozygous mice show little change in bleeding time (Table 1).

Role of the P2Y12 receptor in shear-induced platelet activation
Pharmacologic blockade of the P2Y12 receptor in physiologic conditions of arterial flow revealed that this receptor is essential for platelet aggregation under shear conditions. Blood from a patient with defective P2Y12 receptors, or normal blood treated with the reversible P2Y12 receptor antagonist AR-C69931MX, exhibited small and loosely packed thrombi, whereas normal individuals formed large, densely packed thrombi in physiologic flow experiments (42). P2Y12 antagonism also decreased shear-induced platelet aggregation; however, greater inhibition was achieved by antagonism of both the P2Y12 and P2Y1 receptors (43). P2Y12 antagonism has also been shown to decrease P-selectin expression and microparticle formation that is initiated by platelet interactions with vWF (44). Thus, the P2Y12 receptor also plays a role in the formation of platelet aggregates under shear conditions and contributes to thrombus formation on surfaces coated with either collagen or vWF.

Clinical implications of P2Y12 blockade
Consistent with the central role of the P2Y12 receptor in thrombosis, P2Y12 receptor antagonists reduce occlusive thrombosis in animal models. Clinical studies using clopidogrel demonstrate a significantly reduced risk of peripheral artery disease, myocardial infarction, ischemic stroke, or vascular death, in comparison with aspirin therapy (8). Combination therapy with both clopidogrel and aspirin has been shown by the CURE (Clopidogrel in Unstable Angina to Prevent Recurrent Ischemic Events) study to result in enhanced beneficial effects, and this has led to FDA approval of clopidogrel for the treatment of some acute coronary syndromes (45). An ongoing trial for the Management of Atherothrombosis with Clopidogrel in High-Risk Patients with Recent Transient Ischemic Attack or Ischemic Stroke (MATCH) is expected to test the effectiveness and safety of combined clopidogrel plus aspirin therapy versus clopidogrel only in patients who have experienced a transient ischemic attack or ischemic stroke (46). The results of this trial will further clarify the efficacy of combination therapy. Despite its beneficial effects, clopidogrel has been shown to cause the development of the immune-mediated syndrome thrombotic thrombocytopenic purpura (47).
form is required before the onset of drug action is achieved, and an interaction with the cholesterol-lowering drug atorvastatin, which also requires hepatic metabolism by cytochrome P450 3A4, has been identified (48, 49). Post hoc analysis of clinical studies where patients received both clopidogrel and atorvastatin found beneficial effects of clopidogrel in both the presence and the absence of atorvastatin, indicating that the interaction between the two pharmacologic agents does not alter the clinical effect of clopidogrel (50). Another study also found no difference in the clinical outcome of patients taking clopidogrel with atorvastatin or with other statins (51). The clopidogrel metabolite irreversibly blocks P2Y12 receptor function for the lifespan of the platelet. Hence, prior to surgery, clearance of clopidogrel-treated platelets is necessary to prevent bleeding complications. Thus, new research efforts are aimed at discovering faster-acting, reversible P2Y12 receptor antagonists that would allow more control over antiplatelet treatments.

**In vivo analysis of thrombus growth and stability**

With the established role of the P2Y12 receptor as the central point of thrombus formation, in vivo analysis will clarify the mechanism by which the P2Y12 receptor contributes to thrombus growth and stability. Multiple studies have begun to characterize the effects of either P2Y12 antagonism or P2Y12 knockout on the formation of a thrombus using various models of vessel injury. A recent report by van Gestel et al. (15) investigated the effect of the P2Y12 receptor antagonists AR-C69931MX and clopidogrel on in vivo thrombus growth and stability (Table 1). Mechanical injury of mesenteric arteries in rabbits treated with AR-C69931MX or clopidogrel resulted in decreased thrombus height (20% reduction) within the vessel but no change in the bleeding time of the vessel (15). P2Y12 receptor blockade significantly reduced the total duration of embolization with fewer and smaller emboli being produced, and it also reduced the size of the initial thrombus without affecting its stability (15). Thrombin generation was decreased in the AR-C69931MX–treated mice, suggesting that the P2Y12 receptor contributes to the procoagulant response (15). In clopidogrel-treated mice, thrombosis scores were significantly reduced compared with those in controls (52). In addition, there was a delay in the time of initial thrombus formation in the clopidogrel-treated mice (52). Treatment with AR-C69931MX also decreased the reocclusion rate and improved myocardial tissue perfusion in a canine model of coronary electrolytic injury (53). Andre et al. (16) used several approaches to explore the effects of P2Y12 receptor deficiency on thrombus formation. FeCl3–induced vessel wall injury of the mouse mesenteric artery resulted in occlusion of the wild-type mouse vessel; however, the P2Y12–deficient mouse vessel remained unoccluded in eight out of nine mice (16). The appearance of the first thrombus was delayed and only small “unstable” thrombi formed in P2Y12–/– mice (16). There was increased embolization from the thrombus of P2Y12-deficient mice (16). Contrary to the observations of van Gestel et al. (15), more embolization occurred in the P2Y12–/– mice compared with wild-type or heterozygous mice. The differences between these studies are likely due to differences in both species (rabbit versus mouse) and experimental parameters (mechanical injury versus FeCl3 injury) and represent the beginning of our understanding of the effects of P2Y12 antagonism on thrombus formation in vivo.

Data on the role of the P2Y12 receptor in platelet activation and thrombus formation suggest that this receptor is important for the potentiation of many platelet responses and for the formation of a stable hemostatic plug. The tools available for the analysis of P2Y12 receptor function have facilitated characterization of the implications of P2Y12 antagonism, though the studies performed thus far also raise more questions. The differences in embolization between homozygous and heterozygous mice observed by Andre et al. (16) raise several important questions. Would complete blockade of the P2Y12 receptor be a better therapeutic goal, and would it be more beneficial than clopidogrel because of abolished receptor function? Why do results differ depending on the use of a P2Y12 receptor antagonist as opposed to the use of P2Y12 receptor–knockout mice? Is the reason for the increased bleeding times observed in patients and mice lacking the P2Y12 receptor due to unstable thrombus formation or the small size of the thrombus? What are the reasons for differences in embolization in the studies thus far? Answers to these questions are important because, ultimately, definition of the central role of the P2Y12 receptor would translate into the development of a more effective antithrombotic agent, and the answers would affect treatment modalities. The models for thrombus formation have provided varying results regarding the effect of P2Y12 antagonism and/or knockout on thrombus size and embolization. The method of vessel injury, vessel size, and species differences must be considered in the interpretation of the effects of P2Y12 antagonism on thrombus formation. Of course, it is the clinical data that will determine the effectiveness of such therapies, but it is the models of thrombosis that will guide the efforts toward new and improved P2Y12 antagonists. Thus, the large amount of data obtained using clopidogrel treatment, and the studies of van Gestel et al. (15) and Andre et al. (16), may herald the beginning of a new era in antithrombotics.

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