Why Thrombin PAR1 Receptors Are Important to the Cardiac Surgical Patient

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Abstract: Targeting of the high-affinity thrombin receptor protease-activated receptor-1 (PAR1) on platelets represents an exciting strategy to curb the pro-thrombotic complications of cardiac surgery without interfering with the hemostatic benefits of thrombin in the coagulation cascade. The first dedicated PAR1 antagonist to complete safety trials this year has justified expectations, showing no increased risk of bleeding when added to standard anti-platelet therapy but halving major adverse cardiovascular events after percutaneous coronary intervention. In the setting of cardiothoracic surgery with cardiopulmonary bypass, an FDA-approved drug already exists with anti-PAR1 properties: aprotinin has been shown to inhibit thrombin-induced platelet activation in vitro and clinically, through sparing of PAR1 receptor cleavage and activation. Because aprotinin also exerts anti-fibrinolytic effects through blockade of plasmin, this indicates a subtle clinical mechanism of action that is simultaneously anti-thrombotic yet hemostatic. PAR1 antagonists would also be expected to exert anti-inflammatory properties through targeting of PAR1 on endothelium, and this principle has been validated in vitro for aprotinin and newer peptidomimetic antagonists. PAR1 antagonism is likely to remain an active and exciting area of research in cardiac surgery, with newer generations of PAR1 antagonists and recombinant aprotinin variants entering clinical development. Keywords: cardiac surgery, thrombin, receptor, antagonist.

PAR RECEPTORS: SENSORS OF INJURY

Protease-activated receptors (PARs) use a weird and wonderful ligand receptor activation mechanism that allows them to sense changes in the proteolytic milieu. Whereas other receptors recognize ligands carried in solution phase, the PARs receptors carry their ligand (a hexapeptide motif) within their own receptor exodomain. The hexapeptide ligand, however, remains inaccessible to the receptor binding pocket until unveiled by cleavage with a serine protease (1). The newly created N terminus (with the hexapeptide now at the end) folds back into the body of the receptor and docks within the binding pocket (2). From then on, downstream signaling through G proteins and cell activation is similar to other G protein-coupled receptors of the same seven-transmembrane superfamily.

This unique activation mechanism allows PARs to sense the presence of serine proteases in the environment, not just thrombin. Because PARs receptors are found on all cells of the vasculature and the vessel wall, they provide a critical sensing mechanism allowing the body to respond to surgery and cardiopulmonary bypass (CPB; which is known to activate a range of critical serine proteases, including thrombin, kallikrein, plasmin, tryptase, elastase, and others) (3). Three of the four PAR receptors (PAR1, -3, and -4) are cleaved by the serine protease activity of thrombin and can therefore be considered thrombin receptors (4). PAR1 is the high affinity thrombin receptor and PAR4 is the low affinity thrombin receptor on platelets. PAR3 is poorly understood but may be an important thrombin receptor on vascular cells. PAR2 is the odd one out, because it is not a thrombin receptor, being cleaved instead by trypsin, mast cell tryptase, or the ternary coagulation complex of factor Xa-VIIa-TF.

Although PAR1 is recognized for being the high-affinity thrombin receptor, and thus of critical importance to platelet involvement in thrombosis, it should be remembered that other serine proteases, notably trypsin, kallikrein, and low concentrations of plasmin, can also cleave and activate PAR1 (3,5). This is important when considering the effect of serine protease inhibitors in cardiac surgery.

The reason there is so much excitement about the use of thrombin receptor antagonists in cardiac surgery is that they promise to abrogate the pro-thrombotic actions of thrombin on platelets while leaving the coagulation cascade largely untouched—the hope is that thrombotic...
events can be eliminated without causing undue risk of bleeding (6).

**PROMISE OF TARGETING PAR1 IN CARDIAC SURGERY**

Several PAR1 antagonists are in clinical development. The most advanced, which just completed glowing safety trials for use in percutaneous coronary intervention (PCI) and is now in a 10,000 patient phase III trial, is a peptide antagonist based on the hexapeptide ligand sequence (7). This blocking peptide sits in the ligand binding pocket and prevents access to the natural ligand, even when that is generated after proteolytic cleavage of PAR1 with thrombin (8). The phase II TRA-PCI safety trial met its primary safety endpoint, showing no increase in thrombolysis in myocardial infarction (TIMI) bleeding when added to standard anti-platelet care, but showing a 46% reduction in major adverse clinical events (7). Figure 1 shows how PAR1 antagonists can block thrombotic complications by preventing platelet activation caused by thrombin, whereas they do not interfere with the hemostatic properties of thrombin in the coagulation cascade.

Although specific PAR1 antagonists have stolen the limelight in 2007, the first clinical demonstration of PAR1 antagonism came in 2004 through the use of aprotinin in cardiothoracic surgery with CPB.

**PAR1 TARGETING BY APROTININ: TEACHING AN OLD DOG NEW TRICKS**

Aprotinin is a broad-spectrum serine protease inhibitor first isolated from cow lung in 1936. It was shown to be a plasmin inhibitor in 1979, and its clinical anti-fibrinolytic properties were co-discovered in 1987 by groups in the United Kingdom and Holland (9,10). From the first studies in cardiothoracic surgery, aprotinin was recognized to preserve platelet function (10). Elegant electron microscopy studies showed that platelet morphology was completely preserved throughout CPB (11). The critical study into the mechanism of platelet protection came in 1998 from a study by Victor Ferraris, which showed excessive bleeding was linked to activation and degranulation of platelets through the high-affinity thrombin receptor PAR1 (12).

Given that thrombin activates PAR1 through a serine protease mechanism and that aprotinin is a serine protease inhibitor, we hypothesized that aprotinin should possess anti-thrombotic properties by preventing thrombin-induced platelet activation. This hypothesis was controversial at the time, following the ambiguous results of the IMAGE trial into graft patency (13). We first studied the effect of aprotinin on washed human platelets and were able to show a dose-dependent inhibition of thrombin-induced platelet aggregation (14). This was achieved at clinically relevant concentrations of aprotinin: 42.6 ± 21.6% inhibition at 50 KIU/mL (p = .0047), 61.0 ± 25.2% inhibition at 100 KIU/mL (p = .0001), and 86.6 ± 8.9% inhibition at 160 KIU/mL (p < .0001).

We next examined whether aprotinin could inhibit PAR1 activation clinically (15). This study confirmed that (i) thrombin was generated during passage of blood through the bypass circuit; (ii) platelets were activated by thrombin because of cleavage of PAR1; (iii) high-dose (Hammersmith dose) aprotinin prevented platelet activation through PAR1 without affecting net thrombin generation; and (iv) the mechanism of PAR1 protection was by preventing proteolytic cleavage of PAR1. In vitro, the mechanism is definitively through targeting of thrombin-induced PAR1 activation. Clinically, we cannot rule out the possibility that aprotinin may also target plasmin and kallikrein, both of which can cleave and activate PAR1, in addition to thrombin.

This clinical study therefore revealed a subtle “anti-thrombotic yet hemostatic mechanism” of action for aprotinin when used in cardiothoracic surgery (Figure 1): anti-thrombotic by virtue of preventing thrombin-induced platelet activation and hemostatic by virtue of anti-fibrinolytic targeting of plasmin. Thus, like the more modern peptidomimetic PAR1 antagonists, this opportunistic PAR1 antagonist is able to exert anti-thrombotic properties without increasing the risk of bleeding. Better still, because of its additional targeting of plasmin in the fibrinolytic pathway, aprotinin simultaneously delivers anti-thrombotic and hemostatic properties. This is an exceptionally useful pharmacologic profile for a compound used primarily as a hemostatic agent in cardiothoracic surgery.
Similar anti-thrombotic yet hemostatic properties of aprotinin have been observed in animal models of thrombosis and clinically in off-pump surgery (16,17). Meta-analyses of the randomized trials have borne out that aprotinin does not add risk to graft patency but significantly lowers the risk of stroke (18). A possible mechanism contributing to stroke protection is through reduced perioperative platelet activation by thrombin (19). Another contributory mechanism would be through reduced thrombin activation of endothelium, which is expected to yield anti-inflammatory and anti-thrombotic drug effects (20).

CONCLUSIONS

Clinical phase II trials in 2007 seem to have borne out anticipated anti-thrombotic benefits of PAR1 antagonism not linked to an increased risk of bleeding. The first clinical demonstration of PAR1 antagonism, however, came from earlier work using the anti-fibrinolytic agent aprotinin. This possesses PAR1 antagonistic properties by virtue of blocking proteolytic activation of PAR1 by thrombin. It is anticipated that PAR1 antagonism will remain an active field for further development in cardiothoracic surgery with CPB, because it holds the prospect of reducing thrombotic complications without incurring a concomitant bleeding risk or even while realizing a simultaneous anti-fibrinolytic hemostatic benefit.

REFERENCES