Mast Cell–Derived Heparin Proteoglycans As a Model for a Local Antithrombotic

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Abstract

Mast cell–derived heparin proteoglycans (HEP-PG) reside in vascular tissue and serve as a local antithrombotic. Heparin, as used clinically, is isolated from its protein backbone from porcine and bovine gut mucosa. The isolated heparin is an anticoagulant; however, when bound to a protein carrier the heparin conjugates will become antiplatelet agents by inhibiting collagen-induced platelet aggregation and procoagulant activity, as distinct from soluble heparin. HEP-PG, whether soluble or immobilized to a surface, inhibit platelet deposition on the collagen surface in flowing blood. Mimics of HEP-PG, can be tailored to molecules carrying both antiplatelet and anticoagulant (APAC) properties. These molecules can target the vascular injury site and take residence there. Inhibition of thrombus growth using these APACs under these conditions has been demonstrated in several animal models. Although efficacious for antiplatelet and anticoagulant effects, the bleeding time is shorter with APACs than with unfractionated heparin, suggestive of beneficial efficacy/safety ratio. These strategies may be utilized in drug development, where many vascular injury-related problems can be tackled locally.

Keywords
► heparin
► heparin proteoglycan
► antiplatelet
► anticoagulant

Mast Cells As the Origin of Heparin

Heparin, as used clinically, is one specific member of the glycosaminoglycan (GAG) family, originates from intestinal mucosa and provides a protective anti-inflammatory, anti-microbial, and antithrombotic surface against outer invaders. Heparin was originally derived from liver as a phosphatide, which turned out to possess anticoagulant activity.1 Heparin has served for more than 75 years as a source material for modern anticoagulation. The birth of heparin in the United States by the research of Jay MacLean and William Henry Howell has offered a unique and long-lasting clinical tool to manage and prevent thrombosis.1 Heparin is the preferred anticoagulant to prevent contact activation during certain procedures, such as heart and vascular surgery and plasma exchange. The structural studies for the first clinical use of heparin were performed in 1935 by a Finnish doctor, Erik Jorpes, who had moved in 1919 from Finland to Stockholm Karolinska Institute.2 An anecdote of his fanaticism for this material is written in his biography by Runar Beckman: "When his wife was pregnant and the delivery was near some problems arose. Jorpes was working late in the evenings in the laboratory, and when a friend asked him to come home to help his wife he responded that babies are born all the time but heparin is born only once.”3

The unfractionated heparin (UFH) is isolated from porcine or bovine intestinal mucosa and is used in pharmaceutical manufacturing processes as active product ingredient. It is further chemically or enzymatically degraded to lower molecular weight species, which has been the mainstay of acute anticoagulation needs for several decades.
At the time of writing this article, Seminars in Thrombosis & Hemostasis (STH) contains, according to PubMed, as many as 592 contributions where heparin is mentioned. The oldest entry dates back to 1975, by Gallus et al, “Thrombolysis with a Combination of Small Doses of Streptokinase and Full Doses of Heparin”). During the years, many insights into the heparin molecule, its structure and function, as well as laboratory measures and clinical management have been provided by the journal. The history of heparin and STH has shared parallel and complementary routes. The low-molecular-weight heparin, its subcutaneous administration, and even the ultralow molecular species and the antithrombin-binding pentasaccharide and entities without anticoagulant activities have been vividly discussed during the past 40 years. Therefore, it is a great privilege for us to write this article and propose the parent heparin proteoglycan as a model of future drug development.

The GAG surface is typical of all vasculature and organs. The glyocalyx delineates endothelial and platelet surfaces and protects them from simple contact activation. This natural barrier needs to be disrupted in order to initiate coagulation activity, which according to its physiology localizes and limits the action to the site of injury. Under severe coagulation disorders, such as disseminated intravascular coagulation and thrombotic thrombocytopenic purpura, this protective function is lost or altered and generalized thrombosis occurs. For this purpose, the endothelial glyocalyx and the heparan sulfate provide the scaffold to support the potent anticoagulant activity, which according to its physiology localizes and fosters the site of injury while limiting the action to the site of injury. Under severe coagulation disorders, such as disseminated intravascular coagulation and thrombotic thrombocytopenic purpura, this protective function is lost or altered and generalized thrombosis occurs. For this purpose, the endothelial glyocalyx and the heparan sulfate provide the scaffold to support the potent anticoagulant activity, which according to its physiology localizes and fosters the site of injury while limiting the action to the site of injury.

Mast cells are an integral component of the inflammatory triad, together with neutrophils and lymphocytes. Mast cells reside in tissues, in the peritoneal cavity, in the lungs and vasculature, where they in particular surround the outer surface of vessels, in the adventitial layers. This is the very same site where the strongest tissue factor activity is expressed. The deeper the damage, the more coagulation activity occurs within a given vessel. Thus, the regulatory steps for this activation need to be adopted accordingly. The tissue-associated mast cells upon their activation release heparin and chondroitin proteoglycans (HEP-PG and CS-PG, respectively). HEP-PG are large, on average, 750 kDa structures, containing over 10 heparin chains, of various lengths (likely 20–75 kDa) bound to serine–glycine backbone. This heparin structure also supports the activity of enzymes, such as chymase and tryptase, which both exert anticoagulant properties, such as inhibition of thrombin. Moreover, mast cells contain plasminogen activators, being simultaneously devoid of their regulators of plasminogen-activator inhibitors (PAI-1 and PAI-2), as opposed to all other known cell types.

**HEP-PG Are Strong Inhibitors of Platelet Aggregation and Deposition to Collagen**

We isolated serosal mast cells and noted that when activated these release the HEP-PG, which then exhibits a strong inhibitory action against platelet aggregation induced by collagen. UFH has been used as the control for HEP-PG in all these experiments. Also, under high shear rate blood flow over collagen, the platelet deposition is significantly reduced. HEP-PG attenuate platelet–collagen interactions under blood flow via von Willebrand factor (VWF) and platelet glycoprotein (GP)IIb/IIIa-mediated mechanisms. When chondroitinase-treated HEP-PG was used, the inhibitory capacity was maintained, whereas heparinase eliminated the platelet inhibitory reactions of HEP-PG. HEP-PG also arrests platelet thrombus growth at the immobilized collagen surface in vitro and vascular injury sites in vivo. The inhibition of platelet deposition and thrombus growth was evidenced both in solution when spiked in whole blood and upon immobilization on the collagen surface. Indeed, when immobilized directly to the collagen surface, platelet deposition was diminished. In these experiments UFH was significantly more potent anticoagulant via antithrombin than HEP-PG, whereas HEP-PG exerted its anticoagulant action via heparin cofactor II (Coenraad Hemker, MD, PhD, and Suzette Beguin, MD, PhD, unpublished data, 1998). In the in vivo experiments, the locally administered HEP-PG at the site of rat femoral arterial anastomosis maintained patency while saline and UFH failed to do so.

The concept of mast cell-derived tissue regulation of coagulation was studied in human bleeding time (BT) wounds, where allergen-induced challenge in the BT wound prolonged the BT over the period of follow-up. The prolongation of BT started at 60 minutes and was maximal at 120 minutes after the allergen challenge in comparison with the histamine control. Also, lesions of cutaneous mastocytoma patients when subjected to BT wound expressed prolonged BT from the injury site.

In a rodent study of vascular anastomosis, the local administration of UFH versus HEP-PG clearly showed that the HEP-PG could maintain the vessel patency, and the 111-Indium–labeled–infused platelets deposited less to the anastomosis surface with the exposed collagen and vascular matrix in the presence of HEP-PG than in the presence of UFH. Rat femoral arteries were cross-sectioned and the anastomosis was created in a regular manner. Before closing the last sutures the inner vessel surface was flushed with a small volume of solution with HEP-PG and the blood flow was allowed to enter the vessel. In Fig. 1, the uncovered anastomosis surface can be recognized with the visible stitches, whereas in the controls large thrombi were encountered on the vascular anastomosis sites. The HEP-PG has also been shown to carry significant antiproliferative properties in vascular smooth muscle cells, exceeding that of pharmaceutical heparin species.

**HEP-PG Mimetics APACs As Dual Antiplatelet and Anticoagulants**

We have constructed semisynthetic molecule complexes which can be tailored to contain a desired amount of heparin chains and they express both anticoagulant and antiplatelet actions, especially against platelet aggregation induced by collagen. The platelet inhibition becomes a highly specific feature, which the
soluble unfraccionated and low-molecular-weight heparin species do not possess at clinically relevant or even higher concentrations. In contrast, heparin has in fact been shown also to exert some direct activation on platelets as well as the well-known immunological complications of heparin-induced thrombocytopenia and thrombosis.\(^\text{13-15}\) APACs (antiplatelet and anticoagulant constructs) are conjugates with dual antplatelet and anticoagulant properties. We will report the antithrombotic effects of APACs in vitro, and in rat vascular anastomosis and baboon thrombosis models in vivo.

**Methods**

Agonist-induced (collagen, Nycomed Pharma, Munich Germany; Chrono-Par, Chronolog Corp., Havertown, PA; ristocetin, Helena Biosciences, Gateshead, United Kingdom; ADP, Sigma-Aldrich, St Louis, MO) platelet aggregation was assessed in citrate anticoagulated blood in platelet-rich plasma (PRP) (Aggram, Helena Biosciences). The anticoagulant effects of APAC and UFH were compared using activated partial thromboplastin time (aPTT),\(^\text{16}\) prothrombin time (PT), thrombin time, anti-FXa activity, fibrinogen, FVIII:C, VWF activity and antigen, and calibrated automated thrombogram (CAT),\(^\text{17,18}\) measurements. PT and aPTT were always measured using Thromborel-S and Actin FSL reagents (both from Siemens Inc., Germany). In CAT, platelet-poor plasma (PPP) was supplemented with 5 pM tissue factor (TF) and PRP with 1 pM TF (PPP reagent and PRP reagent, respectively, Stago, France).

The intravenously administered anticoagulant efficacy and safety of APACs, UFH, and vehicle (saline) were even compared in a rat tail BT model.\(^\text{19,20}\) In brief, BT experiments were performed under general anesthesia (2% isoflurane-02) and study compounds were administered and blood was collected via femoral vein catheter. Before dosing, blood was collected in 3.2% sodium citrate for aPTT and PT analysis.

APACs, UFH or the vehicle were injected within 30 seconds. At 5 minutes a blood sample was collected to measure aPTT and control the level of anticoagulation. At 10 minutes the tail was cut at 5 mm from the distal tip with a scalpel and immersed in warm (35 ± 2°C) 0.9% saline. The BT was recorded until the bleeding had been ceased at least for 5 to 10 seconds.

The antithrombotic efficacy of APACs and UFH in comparison with vehicle was also studied in two well established models of acute thrombosis in baboons. In a modified Folts model an extracorporeal AV-shunt was created between the femoral artery and vein in an anesthetized animal.\(^\text{21,22}\) The blood flow was controlled by an external constrictor placed on the artery and the flow was monitored with a probe. The artery was injured from outside by cross-clamping twice for 10 seconds with a Martin needle holder (Hegar-Baumgartner TC Gold 14 cm). All side branches in the proximity of the injury were ligated. The shunt was punctured with a needle (26 G) 1 cm proximal to the vascular access for injecting a bolus (4 mg/mL) of APAC, UFH or saline. The injury was treated for 3 minutes with the study substance before being exposed to blood flow. Immediately after recovering the baseline blood flow, an external constrictor was placed on the injury and flow was reduced to 30 to 100 mL/min (reflecting a stenosis of 90 to 30%). The accumulation of platelets on the stenosed artery was detected by the reduced blood flow and recorded as cyclic flow reductions (CFR). At 5 mL/min the artery was considered occluded and the thrombus was dislodged by releasing the constrictor and flushing with saline. After baseline blood flow was recovered, stenosis was reapplied and the experiment repeated.

In the second baboon model, thrombosis was induced by placing collagen-coated polytetrafluoroethylene grafts (2 cm, 4-mm lumen) into an externalized arteriovenous shunt.\(^\text{23}\) The thrombogenic collagen surface was treated for 10 minutes with APAC or UFH (both at 4 mg/mL). Blood flow was initiated (100 mL/min; a shear rate of 265/s) and the deposition of 111-Indium-labeled platelets and fibrin (accumulation of 125-Iodo-fibrinogen) was quantified for 60 minutes in an anesthetized animal. Efficacy, distribution, and retention on-site of locally administered 64-Cu-labeled APAC or UFH (3 mg/kg) were assessed by positron emission tomography (PET) imaging for 50 hours of partially ligated (two loose sutures 1 cm apart) femoral artery anastomoses in rats.

**APAC Inhibits Collagen-Induced Platelet Aggregation with Simultaneous Anticoagulant Efficacy In Vitro**

APAC inhibited platelet aggregation in human PRP induced by collagen and ristocetin (the latter requiring about a 10-fold higher concentration of APAC in contrast to UFH, but not by
ADP. The difference between the doses to reach at least 70% inhibition of platelet aggregation, varied between 3 to 90 µg/mL in collagen-induced platelet aggregation in citrate anticoagulated PRP between the blood donors (Fig. 2). The mean ED50 dose to inhibit collagen-induced aggregation was approximated to 30 µg/mL (n = 12; independent donors; inhibition 54 ± 24%, mean and SD).

The anticoagulant effects of APAC measured by aPTT and thrombin time were comparable or even somewhat better than UFH, while FVIII, fibrinogen, and VWF variables remained unaltered (results not shown) at an equal heparin concentration in vitro. The anticoagulant action of APAC was neutralized by protamine sulfate to a similar manner as with UFH (data not shown).

In the presence of platelets, APAC molecules dose-dependently impaired thrombin generation by initially prolonging the lag time and time to peak (tpeak), while reducing the peak and endogenous thrombin potential (ETP) both in citrate anticoagulated PPP (supplemented with 5 pM TF and triggered with Ca²⁺) and PRP (supplemented with 1 pM TF and triggered with Ca²⁺). In the representative CAT analysis in PRP (Fig. 3A–C), the baseline (phosphate buffered saline; PBS) lag time was 9 minutes, tpeak 20 minutes, peak 58 nM, and ETP 965 nM. At 0.5 µg/mL, UFH prolonged the lag time 45%, tpeak 48% and reduced the peak 11% in comparison with the baseline. At 0.5 µg/mL, APAC prolonged the lag time by 70%, tpeak by 134%, and reduced the peak by 30% in comparison with the baseline (Fig. 3A). Overall, APAC prolonged the lag time 1.6-fold, tpeak 2.8-fold, and reduced the peak 2.7-fold more than UFH. At 1.0 µg/mL UFH prolonged the lag time 75%, tpeak 159%, and reduced the peak 41%, while in contrast, already at 1.0 µg/mL APAC abolished the thrombin generation (Fig. 3B). At 1.5 µg/mL, UFH prolonged the lag time 79%, while the tpeak and peak were immeasurable (Fig. 3C). ETP was not determined for UFH and APAC due to the significantly delayed thrombin generation at heparin concentrations of 0.5 to 1.5 µg/mL.

APAC in a Rat Tail BT Model and Acute Models of Thrombosis in Primates

APAC variables APAC1 and APAC2 (with double coupling efficacy of heparin compared to APAC1), and the UFH prolonged the aPTT and rat tail BT (n = 5–8) in comparison with the saline-treated animals. aPTT was prolonged at a dose of 0.26 mg/kg by eightfold with APAC1 but only doubled with APAC2 over the baseline (Table 1). At 0.96 mg/kg aPTT prolonged by 10-fold with both APACs while UFH had already reached the maximal time measured (180 seconds). At 1.2 mg/kg, also APAC1 induced the maximal prolongation of aPTT while APAC2 clearly (12-fold) prolonged aPTT, which still could be measured. PT was modestly elevated in all...
In the presence of UFH, the artery repeatedly (5 CFRs/27 min) occluded at a flow rate of 100 mL/min (i.e., stenosed by 30%). In contrast, APAC effectively inhibited thrombus formation until the experiment was interrupted, after the follow-up time of 120 minutes. At the end of the experiment, at time point 180 minutes, the artery was restenosed (30% stenosis) and finally for 10 and 15 minutes, respectively. In the control experiments with UFH and saline repetitive CFRs ensued (results not shown).

In the baboon thrombosis model on the extracorporeal collagen graft, APAC reduced platelet deposition on collagen by 34 ± 13% (n = 4, p = 0.01) in comparison with UFH. The distal thrombus propagation was also diminished by 63 ± 11% (n = 4, p = 0.19) (►Fig. 5). Results with UFH were similar to untreated control values (n = 21). Fibrin accumulation was reduced by APAC (45 ± 14%, n = 4), but not by UFH (1.1 ± 0.1%, n = 4, p = 0.001).

In the rat anastomosis model about 10% APAC attached directly to the vascular application site (►Fig. 6). Compatible with the strong retention potential and slow degradation, PET detected nondegraded APAC at the anastomotic sites in rats for over 50 hours (up to 120 hours), whereas UFH was undetectable already after 24 hours (n = 2). APAC and UFH were cleared via urinary pathway.

### Discussion

APAC molecules show unique dual and localized antiplatelet and anticoagulant action uniformly in several in vitro and in vivo models, including rat tail bleeding time (BT) model, baboon extracorporeal thrombosis model, and rat anastomosis model. APAC is a dual antiplatelet and anticoagulant agent that can inhibit the formation and progression of thrombus in vivo, as well as reduce platelet aggregation in vitro.

**Table 1** Comparison of APAC1 and APAC2 efficacy on hemostasis in a rat tail bleeding time (BT) model

<table>
<thead>
<tr>
<th>Treatment</th>
<th>BT Mean (s) ± SEM</th>
<th>n</th>
<th>aPTT postdose Mean (s) ± SEM</th>
<th>n</th>
</tr>
</thead>
<tbody>
<tr>
<td>Saline</td>
<td>360.00 ± 54.79(^a)</td>
<td>6</td>
<td>15.02 ± 0.26</td>
<td>6</td>
</tr>
<tr>
<td>UFH 0.96</td>
<td>1,949.00 ± 276.96(^b)</td>
<td>6</td>
<td>180.00 ± 0.00</td>
<td>5</td>
</tr>
<tr>
<td>APAC1 0.26</td>
<td>619.17 ± 63.085(^a,b)</td>
<td>6</td>
<td>119.14 ± 37.31</td>
<td>5</td>
</tr>
<tr>
<td>APAC1 0.96</td>
<td>986.29 ± 142.04(^a,b)</td>
<td>7</td>
<td>150.20 ± 17.56</td>
<td>8</td>
</tr>
<tr>
<td>APAC1 1.2</td>
<td>2,467.50 ± 130.58(^b)</td>
<td>6</td>
<td>180.00 ± 0.00</td>
<td>6</td>
</tr>
<tr>
<td>APAC2 0.26</td>
<td>803.17 ± 186.69(^a,b)</td>
<td>6</td>
<td>27.50 ± 3.11</td>
<td>6</td>
</tr>
<tr>
<td>APAC2 0.96</td>
<td>790.67 ± 128.45(^a,b)</td>
<td>6</td>
<td>149.10 ± 15.43</td>
<td>6</td>
</tr>
<tr>
<td>APAC2 1.2</td>
<td>1,704.17 ± 264.26(^b)</td>
<td>6</td>
<td>174.83 ± 5.17</td>
<td>6</td>
</tr>
</tbody>
</table>

Abbreviations: APAC, antiplatelet and anticoagulant; SEM, standard error of mean; UFH, unfractionated heparin.

Note: Saline was used as a negative control and unfractionated heparin (UFH) as the positive control. Samples for activated partial thromboplastin time (aPTT) were collected at 5 minutes after dosing. Detection of bleeding time (BT) was initiated at tail cut 10 minutes after dosing. aPTT was measured up to 180 seconds. The average BT(s) and the aPTT(s) with the standard error of the mean are shown. At 0.96 mg/kg UFH versus antiplatelet and anticoagulant (APAC) 1 and APAC2, p values were 0.004 and 0.008, respectively. APACs showed shorter BTs than UFH despite of the relatively equal aPTT prolongation.

\(^a\)Statistically different from the UFH group, with p ≤ 0.05 following Student t-test.

\(^b\)Statistically different from the vehicle group.
vivo experiments. In addition to inhibition of platelet aggregation by collagen and anticoagulation (prolongation of aPTT and thrombin time) in human PRP and PPP, in the absence of any systemic anticoagulation, APACs were efficient with local administration in vivo in two baboon models of acute arterial thrombosis and collagen-induced platelet deposition and fibrin formation. APACs may be beneficial under several vascular applications where localized thrombosis continues to be a problem despite the use of current antithrombotic medications. These current agents are systemically administered and also pose a bleeding hazard, which may also be lessened by the local drug administration due to a low concentration but prolonged action. We obviously need to demonstrate that there is no significant leakage to the tissue and that the vascular healing continues in a physiological manner. Our preliminary experiments after 2 weeks of local application support intact healing and vascular structures at the site of anastomosis, which has been treated with our APAC compounds.

In vitro CAT experiments in the presence and absence of platelets in plasma continue to indicate the dual anticoagulant effect and inhibition of platelet-dependent procoagulant action of APACs. The anticoagulant effects of APACs were comparable with UFH at similar dose levels in vitro as previously reported. In the presence of platelets, APAC impaired thrombin generation initially by prolonging the lag time and tpeak clearly more than equal doses of UFH. In sharp contrast to UFH, APAC inhibited the collagen-induced platelet aggregation in PRP, but similar to UFH, APAC had no effect on ADP-induced platelet aggregation in PRP.

In vivo BT experiments in rats showed a systemic aPTT prolongation compatible with UFH, but again in deviation from UFH shortened BT, despite the dual anticoagulant and antiplatelet in vitro action. The BT method of rat tail transection, where the tail is immersed in warm saline, is uniformly used to study the anticoagulant compounds regarding efficacy but more importantly safety. In the BT experiments the baseline results were compatible with the previously reported data, and intravenously administered APACs- and UFH-modified hemostasis as shown by the prolongation of the aPTT and BT. APAC1 and APAC2 induced 50% less bleeding and prolonged aPTT at least 15% less in comparison with UFH at the similar dose. APAC1 showed dose-dependent effects on prolonging the BT at the selected dose range while APAC2 with the doubled coupling level of heparin chains induced less bleeding. APACs may limit simultaneously platelet function by inhibiting collagen-induced platelet aggregation and modifying the intrinsic coagulation pathway and anticoagulants in vivo. Importantly, no significant changes were detected in platelet number or other blood cell count during and after these experiments.

The two in vivo experiments in baboons are encouraging regarding the dual antiplatelet and anticoagulant action.

In the modified Folt model of acute thrombosis in baboons, APAC applied locally on the crush-injured and stenosed arterial site effectively and consistently inhibited thrombosis in the artery under the moderate shear stress, with the flow rate of 100 mL/min and stenosis of 30% until the experiment was interrupted after 120 minutes. When APAC was addressed with the short exposure of sequentially increased stenosis from 60 to 90% the patency of the artery was further maintained. In addition, in the extracorporeal collagen graft model, APAC inhibited both thrombosis and fibrin accumulation about by 50% over UFH in the 60-minute experiment in baboons. The APAC potency was further emphasized with the distally propagated thrombus over the effect of heparin. APAC has potential to target itself at the vascular injury site. The local administration and residence of APAC on the femoral artery anastomosis was confirmed with the PET scan in rats where the retention time of copper-labeled APAC was at least twice the time of UFH. This finding refers to the capacity of APACs to target at the injury site and to be maintained there for a significant period of time to inhibit unwanted thrombotic response. As the vascular injury triggers the inflammatory cellular triad of the leukocytes, lymphocytes, and mast cells with their heparin proteoglycans released to the tissue, the local heparin reaction is a native physiological defense mechanism.

Overall, APAC dually inhibited collagen- and ristocetin-induced platelet aggregation in human blood, with anticoagulant properties that were at least comparable with UFH in...
plasma. In the absence of systemic antithrombotic therapy, treatment of a highly thrombogenic vascular and collagen surfaces with APAC arrested in vivo thrombus formation in two models of acute thrombosis in baboons. Locally administered APAC alone, with its dual antiplatelet and anticoagulant effects, may limit the growth of thrombus and prevent thrombo-occlusions in various applications. The encouraging data in the BT model would support even the systemic administration of APAC. However, the local application and administration to thrombotic sites of vascular injury would represent a novel modality to treat arterial thrombosis in association with interventions and risk of acute thrombosis. This model of antithrombotic tissue action would be compatible with nature’s GAGs, including large heparin conjugates derived from mast cells.

Acknowledgments

We thank the following collaborators: Marja Lemponen, HUCH, Helsinki, Finland; Jan Roodt, PhD, and Sep Lamprecht, Bloemfontein, South Africa; Ulla Marzec, MSc, and Stephen Hanson, PhD, formerly from Oregon Health & Sciences University, Portland, OR; Charles Laurent, PhD, and Dany Salvair, IPST, Sherbrook, Canada; and Gijs Van Dedem, PhD, formerly from Organon, Oss, The Netherlands.

References