Overview of Vascular Closure

Vascular closure devices continue to emerge and evolve, offering potential benefits to both patients and practitioners.

BY ZOLTAN G. TURI, MD

Each year since 2003, I have had the privilege of publishing a review of the state of the vascular closure world for Endovascular Today, a sister publication of Cardiac Interventions Today. Over time, these reviews have expanded to include clinical management of vascular closure, old and new technology, complications, a review of some of the more important literature relevant to vascular closure, my clinician's view of the state of the vascular closure industry, and some ruminations on new directions to be expected. As the number of vascular closure devices (VCDs) has expanded, I have used these pages to implement a classification scheme so the various technologies could be placed in perspective. Importantly, understanding where individual devices fall in the classification scheme may allow clinicians to anticipate various properties of these devices even before they gain hands-on experience and hopefully predict the circumstances in which new devices will be successful. Finally, I have had the opportunity to use these reviews as a soapbox to discuss not just vascular closure but also vascular access. Ultimately, the results of vascular closure depend on the quality of vascular access, regardless of whether closure is by manual compression or devices. For an up-to-date review of vascular access, please see the January/February 2008 issue of Cardiac Interventions Today.

What follows is a reprint of this year’s Overview of Vascular Closure from the February 2008 issue of Endovascular Today. I have added some commentary in the medical literature section near the end of this article; a study published this summer has, for the first time, demonstrated that the more scientific approach to vascular access described (and preached) in Endovascular Today for the past several years is associated with reduced complications. The classification system first introduced in this review 2 years ago has now been adopted in an increasing number of settings but is included as Table 1 for those who are unfamiliar. Table 2 has been updated to reflect some additional technology, and some additional comments are included on a new class of devices, described as closure begins with access (CBA) devices.

The past year saw the effect of the recession in endovascular procedures trickle down to VCDs: fewer catheterizations and fewer interventions led to at least a slowing—if not a reversal—of the persistent growth seen in VCDs in the past decade. Despite the economic reali-

TABLE 1. CLASSIFICATION SYSTEM OF CLOSURE DEVICES

Glossary for closure device subclassification:
1. Invasive (I) or noninvasive (N); devices that are placed inside the tissue track are classified as invasive.
2. Active approximation (A) or passive (P); devices are considered active approximators if they mechanically approximate the arteriotomy edges, typically with sutures, clips, staples, or by creating a sandwich that holds the arteriotomy edges together.
3. Clot inducing (C), sealant (S), or neither (0).
4. Permanent (P), temporary (T), or no foreign body (0).

Thus, Angio-Seal would be classified as I-A-(IL)-C-T because it is placed inside the tissue track (I), actively approximates the vessel edges (A), leaves an intraluminal foreign body behind (IL), incorporates collagen for thrombosis (C), and leaves behind a resorbable foreign body (T). Perclose is I-A-(IL)-0-P; StarClose and AngioLink are I-A-(E)-0-P; Mynx is I-P-C-T; and Cardiva Catalyst II is I-P-0-0.
ties of a maturing market, diminishing growth, and increased competition, a number of new devices made their appearance. Several devices had significant changes in platform, and at least one important new technological concept was introduced. As in the previous 6 years, I will review the status of the existing technologies, introduce technologies in the works, discuss some of the practical and theoretical issues affecting vascular closure, and comment on a few of the more important articles in the medical literature. I will take the liberty of using my annual soapbox on safety, discuss VCD complications in general, and revisit the issue of retroperitoneal hemorrhage and VCDs.

**EXISTING TECHNOLOGY**

Angio-Seal (St. Jude Medical, Inc., St. Paul, MN), a "belts-and-suspenders" device because it incorporates active approximation of the arteriotomy along with a

<table>
<thead>
<tr>
<th>TABLE 2. VASCULAR CLOSURE TECHNOLOGY</th>
<th>Invasive/Noninvasive</th>
<th>Active/Passive Approximation</th>
<th>Intraluminal/Extraluminal</th>
<th>Thrombosing/Sealing</th>
<th>Temporary/Permanent Foreign Body</th>
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<tr>
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<td>Intraluminal</td>
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<td>Temporary</td>
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<td>No</td>
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<tr>
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<tr>
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<td>Thrombosing</td>
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<td>Compression</td>
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<td><strong>Investigational</strong></td>
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<td>*</td>
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<td>No</td>
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<tr>
<td>Femoseal†</td>
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<td><strong>FDA Approved; Limited Release or Not Marketed</strong></td>
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<td>VasoSeal‡</td>
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Invasive devices are listed before noninvasive, active before passive approximators, and intraluminal before extraluminal in each category. AngioLink (Medtronic CardioVascular, Santa Rosa, CA). EpiClose Plus (Cardiodex, Tirat-Hacarmel, Israel). SuperStitch (Sutura, Inc., Fountain Valley, CA). Therus (Therus, Seattle, WA). Manufacturers of the other devices are noted in the text. *Subclassification of the Arstasis device will require additional information but can be considered a hybrid because it does not actively approximate vessel edges but does create a tunnel into the lumen. †Marketed outside the US. ‡Available from the manufacturer.
thrombosing agent in the tissue track, continues to dominate the vascular closure market. It is favored with a short learning curve, a high success rate even in the setting of full anticoagulation, and a modest (but very important) complication rate. It is handicapped by two properties inherent to the technology. First, the anchor placed inside the vessel produces a transiently visible filling defect in the arterial lumen and is occasionally obstructive, either at the puncture site or with embolization. Second, it leaves a mass of collagen inside the tissue track and a suture that extends from the arteriotomy to near the skin surface, providing both a nidus and a wick for potential infection. Repuncture should be done with caution during the first 3 months, although a small published series demonstrated no complications.

Perclose (Abbott Vascular, Santa Clara, CA) remains popular among interventionists who prefer the well-established surgical approach of suturing arteriotomies. It leaves less foreign body inside either the artery or tissue track, but unlike Angio-Seal, it does not resorb. StarClose (Abbott Vascular) deploys a nitinol clip rather than suture, is simpler to use than Perclose, and is designed not to leave behind any intraluminal foreign body. In general classification terms, it is similar to Perclose, featuring active approximation, a permanent foreign body, and no thrombosing agent; thus, it has less of a nidus for infection but more of a predisposition to oozing after the procedure in fully anticoagulated patients. The latter may be exacerbated by the diameter of the StarClose deployment shaft. Both Perclose and StarClose lend themselves well to immediate repuncture. There is no restriction on reaccess after Perclose; the evidence base for repuncture after StarClose is modest but has worked well in our experience.

The Boomerang ClosureWire (Cardiva Medical, Mountain View, CA) has a unique niche in vascular closure. Unlike Angio-Seal, Perclose, or StarClose, it is a passive approximator, relying on a nitinol disk inside the artery, with a spring mechanism to maintain traction at the arteriotomy inside the vessel until hemostasis occurs. A theoretical drawback is the need to withdraw the relatively low-profile collapsed assembly through the freshly formed plug, requiring additional compression. Its appeal includes the lack of any foreign body left behind (reducing the risk of infection), ability to repuncture with the same considerations as if manual compression had been used, and deployment through the original procedural sheath. A new version, the Cardiva Catalyst II, is designed to provide facilitated thrombosis in the tissue track by exposing two agents on the shaft of the device to stimulate coagulation, platelet adhesion, and platelet aggregation when tension is applied to the disk inside the vessel. As with other passive approximators, the

Figure 1. Anatomic features for femoral arterial puncture. The bottom of the femoral head (A), center line of the femoral head (B), and the approximate location of the inguinal ligament estimated from a line drawn between the anterior superior iliac crest and symphysis pubis (C) are all visible on plain fluoroscopy before puncture. The actual location of the inguinal ligament (D) can be more accurately assessed with angiography showing the point of lowest excursion of the inferior epigastric artery (E). The inguinal crease (F) is an overutilized and potentially misleading landmark. The ideal target for puncture (yellow oval) is a point below the center line of the femoral head.
litmus test for this device will be its success and complication rate in the setting of the vigorous anticoagulation environment of interventional cases. A more extensive list of devices is included in Table 2.

NEW TECHNOLOGY
The Thresholds for Successful New VCDs

Most laboratories cannot afford the shelf space or inventory management issues raised by stocking more than two or three closure devices. A successful new device in the increasingly crowded VCD marketplace has to meet one or more of the following standards:

- A high enough success rate in both diagnostic and interventional procedures to be the primary, go-to device in the lab
- Favorable features (ease of use, short learning curve, slick deployment mechanism)
- A niche that is perceived valuable
  - Perceived low risk of associated infection
  - Favorable features for use in peripheral vascular disease or puncture outside the common femoral artery
  - Favorable features for use in nonfemoral access

- Manufacturing costs that allow a sustainable profit

The last item, manufacturing costs, may seem tangential to the other considerations, but I suspect this has been the primary cause of some otherwise novel technologies never making it to market. The original VCD, VasoSeal (Datascope Corp., Montvale, NJ), consisted of a few molded plastic parts and one or two collagen plugs. More complex technologies, with finely milled pieces made of expensive metals and multiple moving parts can be prohibitively expensive to manufacture. Device failure, not just failure to achieve hemostasis but failure to function perfectly, is not acceptable to clinicians, patients, or their lawyers; thus, the technical demands in this crowded intellectual property space require substantial creativity.

Return of the Unanchored Plugs

After the demise of both the Duett (Vascular Solutions, Inc., Minneapolis, MN) and VasoSeal (both available from the manufacturer but no longer actively marketed), it appeared that the potential drawbacks of passive closure were proving to be a significant factor in VCD success rates and acceptability. Although both devices had secular issues (VasoSeal had a high failure rate, particularly in fully anticoagulated patients, whereas...
the Duett was associated with occasional intra-arterial injection, sometimes with catastrophic results), the lack of active approximation was perceived to be a drawback for use in the interventional environment. Failure of devices in anticoagulated patients is at best messy, requiring prolonged compression with or without adjunctive use of other devices and is associated with a significant complication rate. The greater success rates of active approximators, such as Angio-Seal and Perclose, relegated the unanchored plugs to small shares in the VCD market.

Thus, it is something of a surprise that the most prominent new VCD marketed in 2007 and the next important device likely to be released are both unanchored plugs. Both devices, the Mynx (AccessClosure, Mountain View, CA) and ExoSeal (Cordis Corporation, Warren, NJ), also share several other characteristics: they utilize biopolymers that are sealing rather than thrombosing agents, both deploy through the existing vascular sheath, and both feature streamlined, short learning curve delivery mechanisms. The Mynx (polyethylene glycol) is being actively marketed, and the ExoSeal (polyglycolic acid) has finished its pivotal trial but is not yet FDA approved. Although both devices appear to have high success rates, yet to be determined are the failure rates in the real-world interventional environment and how well sealing agents stack up against thrombosing agents (ie, biopolymers vs collagen) with regard to tissue track oozing in fully anticoagulated patients.

Given the low single-digit failure rates that operators expect with closure devices in interventional cases, the challenge for these unanchored plugs will be to match that standard in the full anticoagulation/antiplatelet agent environment. If they do, these devices will benefit from their ease of use; if they do not, they will be relegated to that second tier reserved for VCDs used primarily for diagnostic catheterizations.

**TECHNOLOGY IN THE WORKS**

A new class of devices has entered the VCD world, best described as closure begins with access, or CBA. This approach should be distinguished from “preclosure,” typically the deployment of Perclose at the time of initial access and before upsizing the sheath from 6 F or so to very large sizes (up to 24 F in some cases). Preclosure has been around for at least a decade and has had considerable success in settings such as percutaneous stent graft placement for abdominal aortic aneurysms. Now, two true CBA devices have appeared. The FISH (Femoral Introducer Sheath and Hemostasis) device (Morris Innovative Research, Bloomington, IN) uses small intestinal submucosa wrapped around the access sheath, which is deployed as the sheath is withdrawn at the end of the case. This device is FDA approved. Considerable interest has been provoked by the initial presentation of data on the Arstasis device (Modesitt, San Carlos, CA). This technology creates a shallow-angle access track through the femoral arterial wall at the time of puncture to create a self-sealing mechanism as the sheath is withdrawn at the end of the case.
of the case (the developer of the technology prefers the term tunnel to dissection plane to distinguish the Arstasis access mechanism from the types of pathological and uncontrolled dissection with which clinicians associate the term). The Arstasis concept leaves no foreign body behind. A small, first-in-man pilot study presented at the TCT 2007 meeting was reasonably successful. Several important issues need to be addressed before it will be possible to meaningfully comment on the long-term future of the Arstasis concept:

- The applicability of devices based on this technology to interventional cases
- The applicability to femoral arteries with atherosclerosis and particularly calcification

In summary, the Arstasis concept offers a promising approach to arterial access in intervention, but further development is required to address critical issues before it can be widely adopted.

### TABLE 3. AN ALGORITHM FOR PREVENTING AND MANAGING RETROPERITONEAL HEMORRHAGE

1. Access using the iterative fluoroscopic technique described at length in our previous annual articles, with emphasis on puncture over the lower half of the femoral head (Figure 1).

2. Femoral angiography before anticoagulation will reveal a high stick and allow the operator to postpone elective interventions. The postponement is an inconvenience to patients and families and problematic to catheterization lab administrators and insurers. Nevertheless, delaying a procedure for 24 hours is far preferable to RPH and its associated 5% mortality rate. RPH in unanticoagulated patients is rare.

3. In the setting of high stick in a patient who is already anticoagulated, use of VCDs may have significant additive risk. In the study by Ellis and colleagues, the odds ratio was 2.8:1, and although it was apparent only for Angio-Seal, it may well be a class effect. In an earlier study from the same institution, RPH despite IIb/IIIa use occurred in <0.2% of patients when manual compression was used and >1% when VCDs were used. I have interpreted that study to suggest that not only is VCD use a potential culprit but also anticoagulation, because VCDs are deployed in fully anticoagulated patients on the catheterization table, whereas manual compression is performed with the activated clotting times at or near normal levels.

4. Institutions should consider a set algorithm for treating possible RPH after catheterization (Figure 4). Hypotension should always raise the possibility of RPH. The potential for missing the diagnosis is simply too high, and mortality continues to occur too frequently to leave this to ad hoc diagnosis and treatment. Figure 5 shows the relative sensitivity of various findings for RPH after catheterization, based on data from Farouque and colleagues. Note that the most sensitive and specific marker of hemorrhage is, of course, anemia, but because of the equilibration time required for the blood count to fall to diagnostic levels, the confounding effect of dilution typical after catheterization, and the time frame in which diagnosis needs to be made, it is best to make the diagnosis before the blood count becomes diagnostic. Note that the algorithm in Figure 4 features two pathways: one for relatively stable patients and one for patients in obvious trouble—those who are in shock and have not responded to the usual measures, such as fluid bolusing. The CT scanner, although providing the best tool for diagnosis of a stable patient, may not be a suitable location for one who is exsanguinating. We believe that when the facilities and staff are available, unstable patients should be brought emergently to the catheterization lab, and a catheterizer trained in peripheral intervention should obtain contralateral access and be prepared to tamponade a bleeding external iliac artery. If this does not succeed in resolving the hemorrhage after balloon release, a covered stent can be considered depending on anatomy.

5. A sine qua non while these maneuvers are being performed is transfusion at the earliest possible opportunity. Two sources of error frequently confound diagnosis and treatment. An ultrasound at the puncture site may be of little use in making the diagnosis of RPH. Chest pain and ischemia on EKG can be a reflection not just of acute occlusion of the freshly intervened upon coronary artery but can instead reflect a combination of decreased oxygen-carrying capacity and decreased perfusion pressure, thereby causing ischemia in unrevascularized areas of myocardium.
• The potential implications of high or low femoral access
• The ability to use these devices in patients with perivascular fibrosis, such as is seen after multiple femoral access procedures
• The nature of postprocedure healing as compared with manual compression or current VCDs

Both of these devices raise the issue of a need to evaluate the femoral artery before access so as to avoid small or diseased arteries, if indeed these represent limitation to use of these devices. The potential need to enter a healthy segment of the common femoral artery may help speed up an evolution of two approaches I have advocated in Endovascular Today in the past. First, there will be benefit from performance of better and more comprehensive evaluation of the common femoral artery for disease and level of bifurcation before access is obtained. Second, use of fluoroscopic and ultrasound techniques can ensure entry into the ideal target zone (Figure 1) in the common femoral artery rather than in one of the bifurcation vessels or above the inguinal ligament.

VCDs FOR OTHER APPLICATIONS
Several VCDs are being adapted for applications other than vessel closure. A logical consideration for suture technologies has been expansion to closure of patent foramen ovale (PFO). Sutura, Inc. has had a recent first-in-man series with the SuperStitch EL, a modification designed for percutaneous PFO closure. At least one PFO was sealed in 2006 with Perclose, and a spin-off from Abbott (Ovalis, Mountain View, CA) has been developing a percutaneous device for this indication. Cardica Medical (Redwood City, CA), which is developing a VCD, is also developing a PFO closure device, although the nature of their technology is not in the public domain.

THE DARK SIDE
All enthusiasm for VCDs needs to be tempered by the dark side of all medical devices: complications. In the VCD world, this issue is exacerbated by the continuing unresolved issue of the risk/benefit ratio of VCDs versus manual compression. Figure 2 shows the relative risk of VCDs versus manual compression in a number of meta-analyses and propensity analyses. There is tremendous noise in these data and, as discussed in several of our previous reviews in Endovascular Today, the results are muddled by learning curve issues, changing device platforms, changing clinical practices, and the inclusion of devices or generations of devices that have been supplanted by better technology.

Retroperitoneal Hemorrhage Revisited
Nevertheless, it is clear that some complications are additive to manual compression. These include infection (discussed in detail in last year’s review), vascular obstruction, RPH, and possibly nerve entrapment. RPH has been discussed in previous years in this article, but in lecturing on this subject, I am reminded that awareness of the potential additional risk of deploying VCDs in high sticks has not been adequately disseminated to the interventional community.

The salient factors are as follows: high sticks, those above the inferior epigastric artery’s lowest point of excursion (Figure 1), are associated with an odds ratio as high as 17:1 of RPH. The mechanism has obvious and somewhat more subtle features. The obvious is the potential for free bleeding into the retroperitoneal space once the inguinal ligament has been crossed. The less obvious is the mechanism of failure when a closure device is utilized. Figure 3 shows why a plug (and possibly a stitch, clip, or other element in a closure device approaching through the tissue track) would fail to land on the arterial surface: the presence of layers of tissue, notably the transversus abdominis muscle, obstructs passage down to the artery.

Although still lacking a solid evidence base, several straightforward recommendations for postprocedure management deserve to be emphasized (Table 3). Ultimately, RPH continues to challenge excellent institutions and interventionalists. It is unfortunate, because it remains a cause of mortality in every hospital. In my opinion, if the routine steps in Table 3 are followed, the rate of RPH and its consequences can be decreased substantially, although unfortunately not eliminated.

The FDA Database
Although suffering from grossly incomplete reporting, the FDA Manufacturer and User Facility Device Experience (MAUDE) database remains a treasure trove for assessing the complications associated with technology including VCDs. I reviewed the reports for 2007 available as of February 2008. It is important to point out that a minority of complications are reported, that the details of individual cases are notoriously incomplete, that the
data are replete with noise, and there is some duplication. No Clinical Events Committee adjudicates these reports, and thus assignment of causality is hazardous. Further, different institutions, and for that matter, different vendors have disparate reporting standards.

The five closure devices with a significant footprint in the database (Angio-Seal, Perclose, StarClose, Mynx, and Cardiva Catalyst II) had a total of 1,499 adverse event reports in 2007, including 22 deaths, of which several may not have been device related. Of these 22, 15 were due to bleeding, almost all were due to retroperitoneal hemorrhage, four were due to infection, and three were complications of vascular obstruction. The devices had various propensities for mechanical failure, obstruction of the artery, need for surgical removal, infection, pseudoaneurysm formation, and most importantly, blood loss. It is important to point out that there is no MAUDE database for manual compression, and despite Figure 2, the verdict may never be in on a clear risk-benefit ratio.

**THE MEDICAL LITERATURE**

Five articles deserve particular mention from the past year. First, and particularly gratifying, is preliminary confirmation that the techniques promoted in this article for the past half decade appear to be having a positive impact on complication rates. As an extension of a project by the Northern New England Cardiovascular Study Group, Fitts and colleagues had from the Eastern Maine Medical Center reported on 2,651 patients in which outcomes after fluoroscopy-guided vascular access were compared to those in which no fluoroscopy was used. Even though many of the access techniques described in the February 2008 issue of *Endovascular Today* were not used, the length of stay was shorter, and there were fewer arterial injuries after fluoroscopy-guided access. At the same time, the overall complication rate of vascular access and closure is decreasing, as shown in an analysis of more than 36,000 PCI patients from the Northern New England consortium. In the interval between 2002 and 2006, the rate of major vascular complications decreased from 3.4% to 2%. The extent to which this resulted from better access techniques, better adjunctive sheath and pharmacological management, or better VCDs and better VCD deployment techniques is unknown. For a surgeon’s perspective on VCD complications, including a suggested algorithm for complication management, the latest article on this subject by Eidt and colleagues is enlightening.

A cost-minimization analysis of VCD versus manual compression by Resnic and colleagues suggests potential cost savings with VCD use, despite the cost of these devices, largely based on a lower complication rate with VCDs—a finding that will not apply universally to all hospitals, operators, or types of VCDs. Finally, a carefully conducted propensity analysis of nearly 13,000 patients undergoing diagnostic catheterization and PCI showed statistically significant lowering of complications with VCD use (Figure 2). This study is part of an overall trend suggesting improving VCD results and hopefully reflects the increasing attention being paid to vascular access and closure in general.

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