Paravalvular leak (PVL) occurs when there is backflow around a prosthetic valve. This can occur through a variety of causes. Paravalvular Leak is accompanied by regurgitation and is often a significant problem for patients with bioprosthetic or mechanical heart valves. Often manifesting as heart failure (85% of all presenting symptoms) and hemolysis (13–47% of all presenting symptoms and signs), PVL has a prevalence rate as high as 5–17% of all heart valves.4–6

The mechanism of leaks is not well understood. The alignment between sewing ring and annulus may be incomplete because of significant annular calcification. The tissue around valves can weaken as a result of chronic infection. Even the sutures themselves may not allow significant apposition of the valve with the annulus. All of these factors can lead to significant PVL.

How can PVL be treated? Unfortunately, repeat surgery portends a worse prognosis, with mortality rates for the first redo, second redo or third redo surgery of 13%, 15%, and 35%, respectively.2 Each repeat operation is less likely to be successful. Therefore there is ample room for percutaneous approaches.

It is important to select the right patients for PVL closure. Prior to beginning a case, it is important to exclude active infection, valve instability, and/or cardiac thrombus.7 Indications for PVL closure include patients with significant regurgitation accompanied by symptoms of congestive heart failure and/or hemolysis. Important contraindications to PVL closure may include presence of active local or systemic infection, active ischemia, mechanical instability of the prosthetic valve, intracardiac thrombus, and patients with a life expectancy due to comorbidities that is less than 6 months.

Paravalvular Leak: Imaging
Successful PVL closure begins with efficient imaging. There are a variety of methods available to diagnose PVL, including transthoracic echocardiography (TTE), transoesophageal echocardiography (TEE), computerised tomography (CT) and magnetic resonance imaging (MRI). Among these, there is no one optimal diagnostic method; each has significant advantages and disadvantages. The important pieces of information to ascertain during imaging are location, size, shape, severity and the number of leaks.

It is important to know the valve type prior to any imaging (often available from the surgical report), and it can also be helpful to know its orientation and the suturing technique that was used. Orientation of the valve varies, but often the preferred mitral orientation for tilting-disc valves is with the major orifice toward the left ventricular free wall as opposed to the septum. For the aortic position, the preferred orientation is with the major orifice to the right posterior aortic wall. Suturing techniques may also vary, ranging from non-everting mattress (with or without sub-annular pledgets), everting mattress (with or without supra-annular pledgets), simple interrupted, figure-of-eight, and continuous/running sutures. The choice of suture technique depends on valve type and surgeon preference.

Echocardiography allows for direct comparison of pre- and intra-procedural results. However, echocardiography is prone to artifact from prosthetic shadowing.7 Aortic PVL can be diagnosed...
and evaluated often by TTE, whereas mitral PVL often requires TEE (although TTE may be useful for original diagnosis). 3D echocardiography adds the ability to determine the path of a leak, which can often take a serpentine course. The authors perform all PVL interventions with TEE guidance (often with 3D characterisation of leak size and course).

CT and MRI add further information. Retrospective ECG-gated reconstruction allows diastolic and systolic characterisation. CT can also provide the accurate imaging angle for intervention and closure. Unfortunately, artifacts secondary to calcification or the valve can blur the leak itself, making it difficult to visualise. CT, in comparison to MRI, has better spatial resolution, however requires contrast dye and involves exposures to more radiation.6

**Location**

A clock face is often used to describe both aortic and mitral PVLs (Figure 1, Figure 2). The three commissures are assigned an hour on the clock face: between left and right coronary sinus is 5 o’clock, between right and non-coronary sinus is 8 o’clock, non-coronary and left coronary sinus is 11 o’clock. This lexicon helps in communication between imager and operator and also helps to monitor leaks pre- and post-closure. Statistically, aortic leaks are most often between 7 and 11 o’clock (46 %) and also between 11 and 3 o’clock (36 %).9

Both clock face and anatomical criteria can be used to describe mitral PVL location. Location is based on the mitral valve annulus and is described as medial, lateral, anterior or posterior. The clock face for the mitral valve starts at the 12 o’clock position between the aortic valve and mitral valve A2, then the 3 o’clock position is the postero medial commissure and interatrial septum and the 6 o’clock position is the posterior annulus midpoint. According to this system, mitral PVL is found often between 10 and 2 o’clock (45 %) and between 6 and 7 o’clock (37 %).6

**Sizing**

Although the course of the leak may be serpentine, with an orifice that is crescentic or oval in shape, some assumptions can be made about size of the PVL. Echocardiography of the vena contracta of the leak can be used to estimate the size, although this method is not perfect. With the advent of 3D imaging, the leak can be measured in multiple directions. CT and MRI may provide more information if an echocardiogram is unclear. The authors do not recommend balloon sizing as it is associated with a risk of balloon rupture as a consequence of sharp edges due to annular calcium. Measurement of the leak will dictate the choice of device for PVL access, and in turn the delivery system guide or sheath size.

**Paravalvular Leak: Access**

Aortic or medial mitral PVLs can be approached by transfemoral access. The authors’ preference is to use a 0.035" wire, often a hydrophilic one (e.g. Terumo Glidewire, Terumo Medical Corporation, Somerset, NJ, USA) inside a 5 Fr diagnostic catheter (JRA or MP). The wire crosses the leak and the catheter follows the wire. This wire is substituted for a stiff 0.035" wire (e.g. Amplatzer Extra Stiff Wire, St. Jude Corporation, Minneapolis MN, USA). The delivery system guide or sheath is then advanced over the stiff wire, which is then removed, and the device is placed in correct position. If more support is needed for the catheter or delivery system to cross the leak, a rail can be made (either by transseptal access and snare or by externalising the wire through transapical access). If the aortomitral curtain is crossed, it is often necessary to protect this from significant stress by covering a bare wire with a catheter at all times. For a medial mitral leak, a JR4 or IM catheter can be helpful.

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**Figure 1: Location and Sizing of Aortic Paravalvular Leaks**

Aortic paravalvular leak in patient after CoreValve transcatheter aortic valve implantation. Here we see one aortic valve in 120 degree view (A) that on biplane view (B) has at least two areas of major leak (blue arrows). A schematic of part B is shown in C, where the leaks are shown in relation to the IAS and the AV. One leak, which is green, is located near the IAS at the 10-11 o’clock position on the clock face, which is also posterior and medial. This would be in the region of the noncoronary cusp of the original AV. The other leak (in orange) is located near 4 o’clock on the clock face, and is mostly lateral and located in the region of the original right coronary cusp. The bottom of panel C shows how the image intensifier would visualise the lesion. An extreme left anterior oblique (LAO) angulation would overlap the orange and green paravalvular leak above each other. However, an right anterior oblique (RAO) angulation would allow both to be seen adjacent to the valve. In D, we see a post-procedure RAO angulation on fluoroscopy, with two devices in each of the major paravalvular leaks on each side of the CoreValve. AV = aortic valve; IAS = interatrial septum.

**Figure 2: Location and Sizing of Mitral Paravalvular Leaks**

Images for patient with two mitral paravalvular leaks (asterisk) at various locations in relation to the Ao, LAA and IAS (A). In the schematic shown in B, one can see that the anatomy is distorted, with the MV being more medial compared with normal anatomy. The two leaks are seen as well. On the clock face (orange disc) with the aortic valve as the 12 o’clock position, one leak is at the 2 o’clock position (one asterisk) and another leak is at the 9 o’clock position (two asterisks), in relation to spatial position, one leak is anterior and medial (one asterisk) and the other leak is lateral (two asterisks). This illustrates the difficulty of using one naming system, especially in distorted valve anatomy after replacement. The lateral leak is shown in C as a large crescentic leak, with measurements of 16 x 6 mm. The anteromedial leak is shown in D, with a measurement of 9 x 3 mm. Ao = aorta; IAS = interatrial septum; LAA = left atrial appendage; MV = mitral valve.
Paravalvular Leak

**Figure 3: Aortic Paravalvular Leak, Single Device**

74-year-old male patient with paravalvular leak in relation to mechanical aortic valve. The leak is an eccentric leak in the region of the non-coronary cusp and measured 10 x 3 mm on echocardiogram. Right femoral arterial access was obtained and a 5 Fr sheath was placed. A 5 Fr Alligator guide catheter and Terumo hydrophilic 0.035" wire were used to cross the leak in a retrograde fashion (A). This catheter was exchanged over an Amplatz Extra Stiff Wire for a 7 Fr Cook Shuttle Sheath, which was placed in the left ventricle (B). A 0.014" Ironman Wire was placed as access protection through the paravalvular leak into the left ventricle. A 10 mm PDA occluder device was attempted but was unsuccessful in closing the defect (C) and was removed. The 0.014" wire stayed in place (D). A 4 Fr 125 cm AM catheter was placed coaxial inside the shuttle sheath to traverse the paravalvular leak over the 0.014" wire and to reestablish the shuttle sheath across the leak (not pictured). Then a 12/3 mm AVP III device was placed across the leak (E). After fluoroscopic and echocardiographic confirmation of minimal leak and good valve function, the device was released (F).

**Figure 4: Aortic Paravalvular Leak, Multiple Leaks**

After having a mechanical aortic valve in 2008, this patient presented 4 years later with symptoms of heart failure and severe regurgitation. A 5 x 11 mm paravalvular leak was noted near the left coronary cusp and 5 x 8 mm paravalvular leak in the area of the non-coronary cusp (A and B). A 5 Fr MP catheter and 0.035" hydrophilic wire was used to cross the leak near the non-coronary cusp. This was then exchanged over an Amplatz ES 0.035" wire (C) for a 10 Fr Cook Shuttle Sheath. This was then used to advance an AVP III 5 x 14 mm device (D), which was then deployed (E). Similar access was obtained through the other femoral artery and a similar technique was used to cross the leak near the left coronary cusp. An AVP IV 5 x 14 mm device was also implanted (F, G, H). Follow-up echocardiographic views at 42 and 117 degrees show the position of the non-coronary cusp (blue arrow) and left coronary cusp (red arrow) devices (I, J). Both aortic and mitral valve leaflets moved well on transoesophageal echocardiography.

Sometimes transseptal access is needed, either for mitral PVLs or difficult aortic PVLs. Any transseptal system should be used, and this should be performed under transoesophageal guidance. The authors recommend an inferior and midway between superior and posterior position for puncture for most leaks, although it is just as important to make sure the transseptal puncture is performed safely as it is to find a specific spot to cross the septum that will allow crossing of the leak. For difficult aortic PVLs, the leak is crossed during retrograde femoral approach and a rail is formed by snaring the wire in the left atrium. For transseptal access, heparin 10,000 units should be administered.

When a mitral PVL cannot be crossed through other methods or if there are mechanical heart valves in both aortic and mitral positions, transapical access can be considered. In addition to echocardiographic/fluoroscopic visualisation to determine the position of the ventricular apex, it is also important to perform concomitant coronary angiography to avoid the coronary arteries. A sheath (often 4 Fr) is delivered and heparin is given. A device is often used to close the entry site (e.g. Amplatzer PDA Occluder, St. Jude Corporation). The authors recommend this as a third and last option, as there is an increased risk of complications from tamponade, hemothorax or puncture of a coronary artery. Follow-up TTE and chest X-ray are highly recommended at 24 hours after procedure/discharge.

**Paravalvular Leaks: Device Selection**

There are only a few devices designed specifically for PVLs, thus other devices have often been used. The ideal device has the appropriate size and shape for the leak and does not interfere with the valve leaflets. Furthermore, it does not interfere with other vital structures, such as the coronary ostia in the case of aortic valves or left ventricular outflow tract in the case of mitral valves. Optimally, only one device is needed.

The device size is dependent on measurements from echocardiogram (TEE and 3D whenever possible). Angiography helps in the case of aortic PVL when this can be seen next to the valve. Some may use external catheter size to approximate leak size, but this is also dependent on calcification and tortuosity, which can cause difficulty in a catheter’s ability to cross the leak. The authors follow this general algorithm: for a small cylindrical leak, an Amplatzer Vascular Plug (AVP) II or PDA Occluder may be best; for an oval or crescentic leak, the AVP III is more ideal; if the leak is small or has significant angulation, an AVP IV is better as it is more flexible.

Recently, the Occlutech PLD (Helsingborg, Sweden) device has obtained CE mark approval. There are two devices, one square and one rectangular, both made of nitinol braided mesh. Waist size is chosen similar to the defect size, and this ranges from 3 to 7 mm with circular waist for the square device (requiring 5–7 Fr sheath) and from 4 x 2 to 12 x 5 mm for the rectangular device (requiring 5–8 Fr sheath).

**Putting it all together: Aortic Paravalvular Leak**

Retrograde transfemoral approach is the optimal strategy for aortic PVL. TEE is used for imaging, with description of the leak on the clock face as described above. Once the leak is crossed with a hydrophilic wire, it is crosssed again with a 5 Fr diagnostic catheter (often JR4, MP or Amplatzer-1). Defect size determines device size, which dictates size of the guide catheter or long sheath to deliver the device. This is exchanged...
over a stiff wire (often Amplatz Extra Stiff) for a delivery system guide catheter or sheath (e.g. Cook Shuttle Sheath, Cook Corporation, Bloomington, IN, USA). The device is then delivered through this delivery system. TEE then assesses changes with the regurgitant jet. Prior to release, it is important to check for absence of coronary ostia coverage and free movement of the valve leaflets.

If there is significant tortuosity or difficulty crossing the leak, more support can be obtained by building a rail. A transseptal rail involves transeptal access with subsequent snaring of the original wire within the left atrium. An apical ventricular rail involves left ventricular puncture and having the wire through the apex. Figures 3 and 4 demonstrate aortic PVL closure.

Subsequent aortic angiography is often necessary to rule out coronary compression and evaluate the valve for regurgitation. The valve should not have an increased gradient and should have free-moving leaflets. If the device was implanted in the area of the non-coronary cusp, special attention should be given to the anterior mitral valve leaflet.

**Putting it all together: Mitral Paravalvular Leak**

After evaluating the leak with TEE, the first approach is to try the simplest approach: retrograde transfemoral approach with a 5 Fr IM or JR4 catheter and a hydrophilic 0.035” guidewire. Once this is across the leak, the wire can be exchanged for a stiff wire and then the delivery system (guide catheter or long sheath). The wire is removed, the device is advanced, and then the device is deployed. By deploying the first disc within the left atrium, it is simpler to visualise on TEE. If there is an issue with support, transeptal access with snare can be used to make a rail. As mentioned above, it is important to protect the aortomitral continuity with a catheter whenever possible.

When the retrograde transfemoral approach is not successful, an antegrade transeptal approach can have some benefits. It is important to cross posteriorly to avoid the aorta and superiorly to have enough catheter room to reach both medial and lateral leaks. A similar approach with a 0.035” hydrophilic wire, 5Fr JR4 or MP catheter, stiff wire exchange, and then exchange for delivery system guide catheter or long sheath is used. If support is still an issue, a transarterial rail is recommended. Another option is to advance the transeptal sheath through the defect. Some centers have used an Agilis system (St. Jude Corporation) if the catheter is unable to reach the defect or if the puncture site was suboptimal. The advantage of this offers is increased steerability; the disadvantage is a larger transeptal puncture and increased cost of the procedure. Figures 5, 6 and 7 show examples of mitral PVLs.

After the device is placed, TEE should show mobile mitral leaflets, open pulmonary veins, and if the leak was anterior, an unobstructed mitral valve. Angiography is insufficient to make this determination. The authors do not recommend release of a device until these requirements are satisfied.

**Device success**

There are many ways to characterise device success, but the device should treat the problem in a manner that allows it to be removed without causing further injury to the patient's heart. If there is satisfactory reduction of the regurgitation jet and free movement of the valve leaflets, the hemoglobin/hematocrit should also be monitored.

**Special situations**

**Multiple leaks**

In cases of multiple leaks, the authors recommend closing the major leak only at first, as if there is significant infection/hemolysis, the offending device can be identified. If multiple devices are placed, perhaps one is not the infectious source and therefore should not be removed. The authors place multiple devices or close multiple leaks if there is uncertain follow-up or with two equally sized large leaks.

One approach for multiple device placement is the same-sheath approach: both devices go through the same sheath one after the other. Device one crosses the leak and is deployed. Next, the wire and delivery catheter are used to cross the leak again, and the second device is advanced and deployed. With this method, only one access is needed. However, the first device needs to be fully released before the second device can be advanced.

Another method is with new access. Contralateral femoral access and device advancement may be sufficient for aortic PVL (Figure 2). In the case of mitral PVL, this requires a transseptal approach and dilating the septal access point to accommodate a larger sheath. The larger sheath should be a sum of the sheaths required for the individual devices (if the two devices need two 6 Fr sheaths, the septum should be crossed with a 12 Fr sheath). Then two (or three wires for three devices) are used to cross the sheath, these wires are exchanged

**Figure 5: Mitral paravalvular leak, one device**
for stiff wires, and the prior large sheath is switched to the multiple delivery systems. The independent devices are then delivered (see Figures 4 and 5).

**Preserving PVL access with device placement**

When the wire crosses the leak only with great difficulty (e.g. tortuous anatomy and/or suboptimal transseptal catheter position), it is also possible to preserve PVL access during device placement with a 0.014” coronary wire (Figure 1). The position of the devices is also used to record hemocardiogram (4). When deploying mitral PVL devices antegrade, it is important to make sure the mitral valve leaflets are not affected by the ventricular side of the device (6) – this requires readjustment and redeployment so the leaflets are not affected (7).

**Complications**

Complications will occur but must be avoided when possible. These include valve interference (3.5–5.0 %), stroke, endocarditis, post-procedural hemolysis, device erosion, emergent cardiac surgery (0.7–2.0 %) and death (1.4–2.0 %). One study showed major adverse events at 30 days (death, myocardial infarction, stroke, major bleeding and emergency surgery) at a rate of 8.7 %. Embolised devices from the aortic position may travel anywhere. Larger devices are less likely to locate cranially and are often found at the iliac bifurcation. The same holds true for devices that embolise from the mitral position, as most are small enough to pass through the left ventricular outflow tract and the aortic valve.

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**Post-procedural hemolysis** is often due to shearing as blood flows through the now smaller orifice at a higher velocity. While this may worsen the clinical condition, this may also be well tolerated and resolves spontaneously after complete endothelialisation. This may take months.

**Long-term survival**

Technical success rate has reported as 77–86 %, and there has been 67–77 % clinical improvement. A study by Ruiz et al. reported long-term survival at 6, 12 and 18 months as 91.9, 89.2 and 86.5 %, respectively. Sorajja et al. found 1–2 year survival after PVL closure of 70–75 % with an estimated 3-year survival rate of 64.5 %.