Transcatheter closure of paravalvular leaks using a paravalvular leak device – a prospective Polish registry

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Abstract

Introduction: Transcatheter paravalvular leak closure (TPVLC) has become an established treatment option but is mostly performed with off-label use of different non-dedicated occluders. The first one specifically designed for TPVLC is the paravalvular leak device (PLD – Occlutech).

Aim: We present initial short-term results of a prospective registry intended to assess the safety and efficacy of TPVLC with PLD.

Material and methods: We screened patients with paravalvular leak (PVL) after surgical valve replacement (SVR). Heart failure symptoms and/or hemolytic anemia were indications for TPVLC. Patients were selected according to PVL anatomy by RT 3D TEE. Only those considered appropriate for closure with a single PLD were enrolled. The procedures were performed via transvascular or transapical access using type W (waist) PLDs only.

Results: Thirty patients with 34 PVLs (18 aortic, 16 mitral) were included. We implanted 35 PLDs with a total device success rate of 94.3% (100% for aortic, 88.2% for mitral). The procedural success rate, encompassing device success without in-hospital complications, was 94.1% (100% for aortic, 93.8% for mitral). During the follow-up period we recorded an increase of hemoglobin concentration (3.9 to 4.1 g/dl), red blood count (11.6 to 12.2 M/mm³) and functional improvement by NYHA class.

Conclusions: Paravalvular leak device type W is a promising TPVLC device, but meticulous preselection of patients based on imaging of PVL anatomy is a prerequisite. A PLD should only be chosen for channels shorter than 5 mm. Size of the device should match the PVL cross-sectional area without any oversizing. Such an approach facilitates high device and procedural success rates.

Key words: percutaneous closure, paravalvular leak, occluder, prosthetic heart valve.

Introduction

Paravalvular leak (PVL) after surgical valve replacement (SVR) may occur in up to 10% of patients, and 2–3% of them will require repair due to heart failure (HF) or hemolytic anemia [1, 2]. The transcatheter PVL closure (TPVLC) was introduced into clinical practice 20 years ago [3]. Transcatheter PVL closure, with proven beneficial effect on HF symptoms and hemolysis [4], has been recently granted a class IIa recommendation by the AHA/ACC [5]. Transcatheter PVL closure was initially performed as an “off-label” indication with ASD/VSD/PDA occluders and later with vascular embolization devices [6–9].

The first TPVLC-dedicated device to receive CE marking was the Paravalvular Leak Device (PLD; Occlutech International AB, Sweden). It has a double-disc design and can have either a narrow or a wide central module (Figure 1). The discs, filled with fabric for improved sealing, are available in square or rectangular shape. The first TPVLC with PLD was reported in 2014 [10]. The present registry is intended to assess the safety and the efficacy of TPVLC using PLD performed after SVR.

Material and methods

Adult patients were scheduled for TPVLC when presenting with hemodynamically significant PVL at a surgically implanted prosthetic valve or a PVL causing hemolytic anemia with the need for at least one packed red blood cell (PRBC) transfusion. Hemodynamic significance...
was defined by (a) HF symptoms (NYHA class II–IV) that could be attributed to paravalvular regurgitation despite optimal pharmacotherapy; (b) presence of ≥ 2+ PVL jet in color Doppler (CD) mapping accompanied by at least one of the following indirect echocardiographic indicators of significant regurgitant flow:
– In case of mitral PVL: (1) systolic flow reversal in at least one of the pulmonary veins, (2) increased calculated pulmonary artery systolic pressure, (3) lack of left atrium (LA) size reduction after mitral valve replacement (MVR) or recurrent and progressive LA dilation in follow-up, (4) forward transprosthetic flow velocity higher than expected with given prosthesis type and size, provided normal function of prosthetic leaflets;
– In case of aortic PVL: (1) holodiastolic flow reversal in proximal part of descending aorta, (2) lack of left ventricle (LV) size reduction after aortic valve replacement (AVR) or recurrent and progressive LV dilation in postoperative course, and (3) forward transprosthetic flow velocity higher than expected with given prosthesis type and size, provided normal function of prosthetic leaflets.

Patients are disqualified from TPVLC in our institution if an indication for classical valve replacement surgery (prosthetic valve instability, need for coronary artery bypass grafting), active infective endocarditis (IE) or unexplained elevation of inflammatory markers (WBC, CRP) is found. In the current analysis, we included only patients treated with one version of PLD with the wide middle module (Waist type). The second available version (Twist) is also a double-disc device, but the middle module is a thin connector. The Waist-type device has self-centering properties, while the Twist type is more useful for narrow channels and multiple parallel narrow channels. Both PLD types are available in square and rectangular shapes in several sizes.

We used either transseptal or transapical (with lateral mini-thoracotomy) access for mitral PVLs and retrograde transarterial access for aortic PVLs. Transvascular procedures were performed under conscious sedation, while transapical ones were performed under general anesthesia [11]. We implanted a single Waist type PLD into each PVL. The size of the device was chosen according to the dimensions of PVL vena contracta by RT 3D TEE. Accordingly, we matched width and length of the PLD middle module (waist) to the minimum (width) and maximum (length) dimensions of the PVL cross-sectional area (CSA) in multi-planar reconstruction – Figure 2.

We refrained from oversizing to avoid any folding of the device and thus to ensure full expansion of both discs. Additionally, we also measured the length of the channel in the plane perpendicular to the prosthesis’ sewing ring (and CSA plane). This dimension reflected the expected distance separating PLD discs after implantation.

Since different TPVLC scenarios, such as repeated implantation during the same procedure as well as repeated procedure after failed implantation, are possible, the following definitions were outlined for this paper:
– Procedure: gaining access + PVL location and crossing + device implantation + delivery system removal + access site protection (suture(s)/closure device);
– Procedure time: from arterial/venous puncture for the transvascular approach or skin incision for the transapical approach until final hemostasis;
– Device implantation: crossing PVL channel with delivery sheath + occluder introduction through delivery sheath + occluder implantation into PVL channel + delivery cable release.

Study endpoints:
– Device success: device implantation with stable device position without interference with prosthetic valve discs/leaflets and with PVL CSA reduction by at least 90%; calculated per number of attempted implantations;
Procedural success: device success + no safety endpoints during hospitalization; procedural success rate was calculated per number of attempted procedures.

Safety endpoints (in-hospital and after 30 days of follow-up): TPVLC-related complications were modified VARC-2 [12] procedure-related complications, additionally including: prosthetic valve disc impingement occurring after plug deployment, PLD embolization, significant exacerbation of hemolysis defined as either procedure-induced hemolytic anemia requiring transfusion of at least 2 PRBC units (once the bleeding-related anemia was excluded) or clinically overt jaundice, and other complications as defined in VARC-2.

NYHA class changes over 30 days of follow-up.

Statistical analysis

Changes were presented as mean and SD values for normal distribution, median and IQ values for non-normal distribution; all calculations were performed with commercially available software (MedCalc v.14.12.0).

The study was approved by the Ethics Committee of the Medical University of Silesia and supported by the STRATEGMED II grant (National Centre for Research and Development, STRATEGMED2/269488/7/NCBR/2015).

Results

Thirty patients presenting with a total of 34 PVLs were included and followed up. Demographic and clinical data are listed in Table I. We performed 18 TPVLC procedures in the aortic location (including 2 PVLs in the same patient closed sequentially during two procedures) and 16 TPVLC procedures in the mitral location (including 2 PVLs in the same patient closed simultaneously via transapical access and a repeated procedure after failure of the first attempt in another patient). The study flowchart is shown in Figure 3.

The mitral location of the PVL was associated with longer procedural time, a higher dose of radiation (Table II) and an considerably lower procedural success rate. In general, such procedures required transseptal puncture, the creation of a loop in the left atrium using the telescopic catheter system. Also, the deployment of the device was more challenging in mitral PVL.

Procedural success was achieved in 32 (94.1%) procedures, more frequently in aortic PVL (18; 100%) than...
in mitral PVL (15/17; 88.2%) cases. There was 1 case of device failure with exacerbation of hemolytic anemia. Table III shows the technical procedural aspects of PLD implantation.

The correlation between the sizes of implanted PLDs and PVLs’ CSA dimensions by RT 3D TEE is presented in Figure 4. Influence of device oversizing on the effect (incomplete vs. total closure) was analyzed separately for the length and the width of PLD – see Figure 5 (incomplete closures are marked with filled circles, total closures with empty ones).

Residual paravalvular flow more often occurred in cases of device oversizing with regards to the length of PVL’s CSA, but the correlation did not reach statistical significance ($p = 0.068$).

There were no major adverse cardiac and cerebro-vascular event (MACCE) during hospitalization and no urgent surgical interventions. In 2 patients small access site pseudoaneurysms developed. They did not fulfill the VARC criteria and were successfully treated by thrombin injection. In one patient there was worsening of hemolytic anemia with jaundice, increase of bilirubin level (from 1.91 mg/dl to 3.2 mg/dl) and lactate dehydroge-

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Table I. Demographic and clinical data

| Parameter                                      | Result              |
|------------------------------------------------|---------------------|
| Number of patients                             | 30                  |
| Age [years]                                    | 63 (IQ 59–70)       |
| Female, n (%)                                  | 19 (63)             |
| Mitral PVL location                            | 13                  |
| Aortic PVL location                            | 17                  |
| NYHA on presentation, n (%)                    |                     |
| I                                              | 0                   |
| II                                             | 7 (23.3)            |
| III                                            | 16 (53.3)           |
| IV                                             | 7 (23.3)            |
| Transfusion – dependent hemolytic anemia, n (%)| 4 (13.2)            |
| HGB [g%]                                       | 11.6 (IQ 10.6–13.4) |
| RBC [M/mm³]                                    | 3.9 (IQ 3.7–4.2)    |
| LVEF [%]                                       | 51.0 (IQ 36.0–57.2) |
| EuroSCORE II                                   | 7.3 (IQ 4.1–11.3)   |

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Table II. Procedural data

| Parameter                     | Aortic PVL                  | Mitral PVL                  |
|-------------------------------|-----------------------------|-----------------------------|
| Procedure time [min]          | median 90, IQR 70–100       | median 125, IQR 95–180      |
|                               | (min. 25, max. 145)         | (min. 50, max. 200)         |
| Radiation dose [mGy]          | median 732.5, IQR 383–1035.5| median 980, IQR 541–1808    |
|                               | (min. 195, max. 1937)       | (min. 267, max. 2220)       |
| Contrast medium volume [ml]   | median 50, IQR 40–67.5      | used only in 5 cases        |
|                               | (min. 20, max. 280)         | (20 ml in 4, 40 ml in 1 patient) |

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Figure 3. Study flowchart

Table III shows the technical procedural aspects of PLD implantation.

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*1 s attempt after failed implantation, **two PLVs addressed in one procedure.

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### Table III. Technical description of individual implantation steps

| PVL location/prosthesis mechanical (m) or biological (b) | Delivery sheath | PLD size | PLD implanted into PVL channel/optimal rotation gained | Release from delivery cable | Device success |
|---------------------------------------------------------|-----------------|----------|-------------------------------------------------------|-----------------------------|----------------|
| Aortic/m                                                  | 8 Fr < 7 Fr     | 10 × 4   | Y/Y                                                   | Uneventful                  | Y (complete closure) |
| Mitral/m                                                  | 9 Fr            | 8 × 4    | Y/N                                                   | Uneventful                  | N (significant residual flow) |
| Mitral/m                                                  | 12 Fr           | 12 × 5   | Y/Y                                                   | Uneventful                  | Y (mild residual leak)   |
| Aortic/b                                                  | 7 Fr            | 8 × 4    | Y/Y                                                   | Uneventful                  | Y (complete closure)    |
| Aortic/m                                                  | 6 Fr > 6 Fr     | 6 × 3    | Y/Y                                                   | Repeated maneuvers necessary | Y (mild residual leak)   |
| Aortic/m                                                  | 6 Fr            | 6 × 3    | Y/N                                                   | Uneventful                  | Y (complete closure)    |
| Aortic/b                                                  | 7 Fr            | 8 × 4    | Y/Y                                                   | Uneventful                  | Y (complete closure)    |
| Aortic/b                                                  | 7 Fr            | 8 × 4    | Y/Y                                                   | Uneventful                  | Y (complete closure)    |
| Mitral/m                                                  | 6 Fr            | 6 × 3    | Y/Y                                                   | Uneventful                  | Y (complete closure)    |
| Mitral/m                                                  | 6 Fr            | 6 × 3    | Y/N                                                   | Uneventful                  | Y (complete closure)    |
| Mitral/m                                                  | 7 Fr            | 8 × 4    | Y/Y                                                   | Uneventful                  | Y (complete closure)    |
| Mitral/m                                                  | 12 Fr           | 12 × 5   | Y/Y                                                   | Uneventful                  | Y (complete closure)    |
| Mitral/m                                                  | 9 Fr > 12 Fr    | 10 × 4   | Y/Y                                                   | Uneventful                  | Y (mild residual leak)   |
| Mitral/b                                                  | 8 Fr            | 6 × 3    | Y/Y                                                   | Uneventful                  | Y (mild residual leak)   |
| Mitral/b                                                  | 12 Fr           | 12 × 5   | Y/Y                                                   | Uneventful                  | N (significant residual flow) |
| Mitral/m                                                  | 9 Fr            | 8 × 4    | Y/N                                                   | Uneventful                  | Y (complete closure)    |
| Aortic/b                                                  | 6 Fr            | 4 × 2    | Y/Y                                                   | Repeated maneuvers necessary | Y (complete closure)    |
| Mitral/m                                                  | 12 Fr           | 12 × 5   | Y/Y                                                   | Uneventful                  | Y (complete closure)    |
| Mitral/m                                                  | 12 Fr           | 12 × 5   | Y/Y                                                   | Uneventful                  | Y (mild residual leak)   |
| Aortic/b                                                  | 6 Fr            | 6 × 3    | Y/Y                                                   | Uneventful                  | Y (complete closure)    |
| Aortic/b                                                  | 6 Fr            | 6 × 3    | Y/Y                                                   | Uneventful                  | Y (complete closure)    |
| Aortic/b                                                  | 6 Fr            | 6 × 3    | Y/Y                                                   | Uneventful                  | Y (complete closure)    |
| Mitral/m                                                  | 9 Fr            | 6 × 3    | Y/Y                                                   | Uneventful                  | Y (mild residual leak)   |
| Mitral/m                                                  | 6 Fr            | 4 × 2    | Y/Y                                                   | Uneventful                  | Y (complete closure)    |
| Mitral/m                                                  | 6 Fr            | 6 × 3    | Y/N                                                   | Uneventful                  | Y (complete closure)    |
| Mitral/m                                                  | 7 Fr            | 8 × 4    | Y/Y                                                   | Uneventful                  | Y (mild residual leak)   |
| Aortic/b                                                  | 8 Fr > 4        | 12 × 5   | Y/Y                                                   | Repeated maneuvers necessary | Y (complete closure)    |
| Mitral/m                                                  | 9 Fr            | 8 × 4    | Y/Y                                                   | Uneventful                  | Y (complete closure)    |
| Mitral/m                                                  | 6 Fr            | 4 × 2    | Y/Y                                                   | Uneventful                  | Y (complete closure)    |
| Mitral/m                                                  | 6 Fr            | 6 × 3    | Y/Y                                                   | Uneventful                  | Y (complete closure)    |
| Mitral/b                                                  | 12 Fr           | 12 × 5   | Y/Y                                                   | Uneventful                  | Y (complete closure)    |
| Mitral/m                                                  | 12 Fr           | 12 × 5   | N/Y                                                   | n/a                         | N (occluder unstable, removed) |

*aFlexor Shuttle Guiding Sheath (COOK Medical) 90 cm, bFlexor Shuttle Guiding Sheath (COOK Medical) 110 cm, cFlexor Shuttle Guiding Sheath (COOK Medical) 110 cm, dAmplatzter TorqVue too short for aortic location accessed from femoral puncture, undersized 8 Fr sheath 30 cm longer used, eAmplatzter TorqVue (St Jude Medical) 80 cm 45°, fAmplatzter TorqVue (St Jude Medical) 80 cm 45° oversized to encompass additional 0.035" control wire, gTelescopic system used for transapical approach, consisting of a longer (23 cm) delivery sheath also used as a PLD loader and wider by 2 Fr, shorter (15 cm) sheath serving as a transapical access, both sheaths normally used for peripheral access, hFlexor Shuttle Guiding Sheath too short – exchanged for an undersized 7 Fr longer (110 cm) sheath, Delivery sheath damaged by the occluder while passing the transversely located PVL channel; exchanged for a new one, the same type, delivery sheath damaged (kinked) in the left atrium; exchanged for a wider one, allowing simultaneous use of a stiff control wire, maintaining proper shape of the sheath inside the LA, iRotation difficulty due to severely angulated delivery sheath, jRotation difficult due to the proximity of surrounding structures.
nase (LDH) activity (from 1214 IU/l to 2294 IU/l). Both decreased spontaneously after 5 days to 1109 IU/l and 2.01 mg/dl, respectively.

Thirty-day follow-up

No additional events or safety points were noted. None of the patients required transfusion, and the hemoglobin (Hb) levels and RBC count increased in comparison to baseline preprocedural values (Hb median of 4.1 (IQ 3.8–4.3) g%, RBC 12.2 (IQ 10.8–13.3)).

Follow-up NYHA class was I in 14 (46.66%) patients, class II in 13 (43.33%) patients, and class III in 3 (10%) patients at 1 month after TPVLC (in patients who underwent a repeated procedure after failure of the first attempt, NYHA class was assessed 30 days after the second one). The difference in NYHA class distribution at baseline and at the 30-day follow-up is presented in Figure 6.

Discussion

Direct comparison of our results with those reported for the so far most widely used AVP III ocluder is difficult because of varying device/procedural success definitions. In one of the earlier papers [13], success, defined as implantation of the ocluder, was achieved in 86%. Later, Cruz-Gonzales et al. [7] achieved the procedural success of 90.9%. It was semi-quantitatively defined as a reduction of paravalvular flow by at least one degree and might correspond to our device success. Another paper on the efficacy of multi-plug TPVLC technique using AVP III reported procedural success, demandingly defined as a reduction of regurgitation to not more than mild, of 76.5% for mitral and 100% for aortic PVLs [9]. Those results were similar to these currently achieved with PLD. Our findings, however, may point to the superiority of PLD in patients carefully selected according to the PVL anatomy.

In the above papers, a multi-plug technique was needed for TPVLC with AVP III devices in the case of large PVLs. Even though efficient, such an approach is also somehow tricky as the increased number of occluding devices may result in their instability. Contrary to that, a PLD ocluder facilitates closure of even large PVLs with a single device. Considering the substantial difference in PVL anatomy suitable for AVP or PLD, both techniques should probably be regarded as complementary rather than competitive, and a head-to-head comparison seems futile. A multi-plug AVP III approach seems reasonable for sealing long, irregularly shaped PVL channels. A single-plug PLD implantation appears superior in more regular, large PVLs with a short channel (shorter than the distance between discs, i.e. 5 mm).

Figure 4. Correlation between the sizes of implanted PLDs and PVLs’ dimensions (length and width of CSA) by RT 3D TEE

Figure 5. Influence of device oversizing on residual flow presence (filled circles) or absence (empty circles)

Figure 6. NYHA class distribution at baseline and at 30-day follow-up
As shown in Table II, most PLD implantations required the use of delivery systems produced by other manufacturers. Even though such provisional solutions enabled the deployment in the majority of cases, we believe that development of a PLD-dedicated system could further improve the success rate. The current lack of such a system may also generate the need for transapical access in some mitral TPVLC cases.

As previously observed, the long-term clinical effect of TPVLC largely depends on the amount of residual flow [14]. Potential mechanisms of its occurrence after PLD implantation, besides the unlikely case of choosing a device with discs too small to cover the PVL, include:

- Occluder deformation caused by elongated PVL – if the length of the channel exceeds 4–5 mm, the discs become spherical. It hinders the proper apposition of the fabric within them to surrounding tissue and excludes proper sealing.
- Squeezing of the waist causing similar disc deformation in cases with a long channel – we observed residual leaks in patients in whom the size of the chosen waist device (PLD type W) even slightly exceeded the dimensions of the PVL.

Technical problems listed in Table II mostly resulted from a lack of TPVLC-dedicated delivery systems and were similar to those occurring with other types of occluders. No PLD-specific complications were identified. Apart from access site-related complications (the same delivery systems as with other occluders), we noted one case of significant post-procedural hemolytic anemia, the risk for which has been described since the beginning of the PVL closure using Paravalvular Leak Device (PLD) [3].

We carried out a pilot study assessing PLD procedural performance and safety/efficacy during short-term follow-up. The changes in anemia parameters and functional class should be interpreted cautiously, because of the small sample size, heterogeneity of the population and changes in medications over the course of follow-up. Long-term follow-up is necessary to confirm the findings.

**Conclusions**

The Waist type of PLD appears to be a useful device for TPVLC in patients after SVR. Meticulous assessment of patients based on imaging of PVL anatomy is a prerequisite. Regardless of PVL CSA length and width, a PLD can be successfully used only for channels shorter than 5 mm. When choosing the best size of the device, one should refrain from any attempts to oversize. In fact, we speculate that slight undersizing may be advisable to avoid any potential deformation of the occluding discs and ensure their proper apposition to surrounding tissue and full sealing. Such use of PLD facilitates high device and procedural success rates. Further elaboration of this technology should probably focus on developing dedicated transvascular delivery systems.

**Conflict of interest**

The authors declare no conflict of interest.

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