Preprocedural imaging for transcatheter mitral valve-in-valve replacement: Planning makes perfect

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1 | INTRODUCTION

Transcatheter mitral valve-in-valve replacement (TMViVR) in degenerated bioprostheses or valvular rings shows promise as an alternative to surgical MVR in selected high-risk patients. However, these procedures are particularly challenging given the complex anatomy of the mitral valve apparatus and the surrounding structures, potentially causing LV outflow tract obstruction (LVOTO). Preprocedural planning with virtual implantation and planimetric estimation of the neo-LVOT at end-systole is crucial for improving procedural results. We present a case of a 84-year-old female patient undergoing TMViVR for degeneration of a mitral valve Mosaic bioprosthesis. Preprocedural planimetric assessment of the neo-LVOT using Materialise Mimics Enlight software allowed for accurate post-procedural results. We also incorporated dynamic assessment of the neo-LVOT during the cardiac cycle and provided immersive visualization using virtual reality imaging.

2 | CASE REPORT

A 84-year-old woman was referred with worsening exercise tolerance and shortness of breath (New York Heart Association [NYHA] class 3–4). In 2011, she underwent surgical mitral and aortic valve replacement (MVR: 29 mm Mosaic [Medtronic]; AVR: 21 mm Mitroflow [Sorin Group Inc.]) for mitral and aortic valve stenosis, followed by dual-chamber pacemaker implantation for postoperative atrioventricular block. In 2017, she underwent transcatheter valve-in-valve aortic valve replacement (TAVR) with an Edwards Sapien XT 20 mm (Edwards Lifesciences)
because of degeneration of the Mitroflow bioprosthesis. In 2019, she presented with recurring dyspnea and progressive degeneration of the mitral valve Mosaic bioprosthesis (mean gradient 5 mmHg at a heart rate of 70 bpm, estimated mitral valve area of 1.6 cm² and severe, eccentric, valvular mitral valve regurgitation) (Video S1A).

DICOM data from 4D electrocardiogram-gated computed tomography (CT) were imported in the Materialise Mimics Enlight software (Mitral Planning Wizard, Materialise) for further assessment. First, using the double-oblique technique, aligning cross-hairs with the mitral plane in a sagittal and coronal view, mitral annular dimensions were obtained: perimeter 76 mm, annular area 450 mm²; largest diameter 26 mm, smallest diameter 23 mm. Second, Sapien 3 26 mm and 29 mm valve frames (Edwards Lifesciences) were virtually positioned within the Mosaic 29 mm in different landing zones (20% atrial/80% ventricular up to 50% atrial/50% ventricular). The smallest left ventricular outflow tract (LVOT) area between the anteroseptal wall of the LV and the simulated valve was segmented, showing a neo-LVOT of 117 mm² (worst case: atrial 20%/ventricular 80% for the 29 mm valve) to 275 mm² (best case: atrial 50%/ventricular 50% for the 26 mm valve). Based on preoperative annular sizing and anticipated neo-LVOT, a 26 mm SAPIEN 3 was selected aiming for a position 30% atrial/70% ventricular (neo-LVOT predicted 176 mm²—Figure 1A).

The patient underwent TMViVR using a standard transseptal approach and transesophageal echocardiography (TOE) guidance.¹ The outflow of the Sapien valve was aligned with the sole radiopaque markers on the struts of the Mosaic valve, to obtain a 70% ventricular position at valve expansion. TOE evaluation immediately post-implantation (Video S1B) showed excellent valve position and function with a mean gradient of 4 mmHg at a heart rate of 80 bpm and absence of mitral regurgitation. A cardiac CT 1 month post-procedure confirmed proper alignment of the Sapien valve with the Mosaic struts, resulting in a neo-LVOT of 171 mm² at end-systole (Figure 1B). Additional measurements at different time points during the cardiac cycle showed a neo-LVOT of 302 mm² at end-diastole and 229 mm² at mid-systole (Figure 1C and Video S2). Using in-house developed software, we were also able to provide an immersive visualization of the neo-LVOT after stent implantation using virtual reality imaging, which can be accessed afterward using a QR code, a smartphone, and a Google Cardboard (Figure S1 and Video S3).

3 | DISCUSSION

Transcatheter mitral valve-in-valve replacement in degenerated bioprostheses or valvular rings shows promise as an alternative to standard surgical MVR in selected high-risk patients. However, these procedures are particularly challenging given the complex anatomy of the mitral valve apparatus and the surrounding structures: Transcatheter mitral valve prostheses may protrude in the LV cavity and cause LV outflow tract obstruction (LVOTO).² Therefore, cardiac CT prior to device implantation is performed to accurately size the mitral annulus, calculate working views, assess access points to the heart chambers, and perform TMViVR simulation to assess the risk for LVOTO. During such simulation, a selection of

![Figure 1](https://example.com/figure1.png)

(A) Preprocedural virtual implantation (30% atrial/70% ventricular with Sapien 3 26 mm). 2D and segmented view with valve in 3-chamber, commissural and short-axis view at end-systole. (B) Postprocedural result with Sapien 3 26 mm in situ indicating excellent correlation with preprocedural virtual implantation. (C) Postprocedural result at end-diastole, mid-systole, and end-systole with Sapien 3 26 mm in a 3-chamber view with a measured neo-LVOT of 302 mm², 229 mm², and 171 mm², respectively
available valve frames and sizes is virtually positioned at different angles and depths of deployment (usually 20% atrial/80% ventricular to 40% atrial/60% ventricular) during the mid- to late-systolic phase, followed by planimetric assessment of the neo-LVOT and estimation of the risk for obstruction. Few studies have evaluated cutoff values for minimal LVOT area that would indicate an increased risk of LVOTO, although Wang et al. found a neo-LVOT area ≤189 mm² to have a high sensitivity and specificity for LVOTO post-procedure (defined as an increase in post-transcatheter mitral valve replacement (TMVR) gradient ≥10 mmHg). There is agreement that patient-specific simulation in TMVR is needed, since LVOTO appears to be the most important predictor of mortality at 30 days. Different preprocedural planning software tools can be used to assess mitral valve anatomy, LV size, interventricular wall thickness, and transcatheter valve size, all of which influence neo-LVOT dimensions. Crucial is that these tools allow for efficient analysis (i.e., become less time consuming) and accurately predict anatomic outcome. This case highlights that rigorous preprocedural planning is critical for appropriate selection of valve type and size, and determining optimal valve positioning depth (in this case 30% atrial/70% ventricular) and resulting neo-LVOT dimensions, while avoiding LVOTO. We think that evaluating neo-LVOT dimensions during the cardiac cycle may further refine our preprocedural assessment both qualitatively and quantitatively. Since the largest part of LV stroke volume is ejected prior to end-systole, further studies have to clarify at what time point during the cardiac cycle the risk for neo-LVOT obstruction is highest. Future studies may further assess dynamic interactions between the valve and host, resulting in more accurate predictions of LVOTO.

4 | CONCLUSION

Transcatheter mitral valve-in-valve replacement in degenerated bioprostheses or valvular rings shows promise as an alternative to surgical MVR in selected high-risk patients, but preprocedural planning, especially planimetric assessment of the neo-LVOT, is crucial. Dynamic assessment of the neo-LVOT may further refine preprocedural planning, but is feasible using dynamic cardiac CT. Immersive assessment through virtual reality imaging may provide additional information since it provides malleable visualization of internal and external anatomy.

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CONFLICTS OF INTEREST
None.

AUTHOR CONTRIBUTION
Alexander Van De Bruaene wrote the manuscript. Stijn De Buck involved in technical support for 3D analysis and virtual reality imaging, and critical review. Peter Verbrugge involved in critical review. Christophe Dubois wrote the manuscript and involved in critical review.

ETHICAL APPROVAL
Ethics approval was not necessary for this case report. The patient’s data and images are de-identified.

CONSENT
Written informed consent was obtained from the patient to publish this report in accordance with the journal’s patient consent policy.

DATA AVAILABILITY STATEMENT
The data that support the findings of this study are available on request from the corresponding author. The data are not publicly available due to privacy or ethical restrictions.

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SUPPORTING INFORMATION
Additional supporting information may be found in the online version of the article at the publisher’s website.

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