Mortality and morbidity among patients with heart failure (HF) remain high, despite advances in therapy.

Heart failure with preserved ejection fraction (HFpEF) is driven by diminished left ventricle relaxation and elevated filling pressures, all of which lead to pulmonary congestion [1, 2]. In these patients, therapeutic options impacting prognosis are limited.

Recently, a novel therapy has been proposed. It is based on creating a communication between both atria using a trans-septal puncture and balloon septostomy. The concept is based on the well-known Lutembacher syndrome, which is defined as a combination of mitral stenosis and atrial septal defect (ASD). The left to right interatrial shunt enables decompression of the left atrium and thus, may improve patient symptoms.

Moreover, observations made in elderly subjects with masked left ventricle restriction, who underwent ASD closure, showed that temporary ASD occlusion with a balloon resulted in significant elevated left atrial pressure [3]. A fenestrated ASD septal occlude has been designed to enable bidirectional flow both in systolic and diastolic impairment [4]. Implantable pressure systems have provided data that left atrial pressure is highly variable over the course of a day and sustained elevations precede clinical events, averaging > 25 mmHg for several days before admission or death [5].

Therefore, a therapy focused on decreasing left atrial filling pressures seems to be promising. Several small studies proved initial safety and efficacy of three different interatrial shunting devices in therapy for patients either with heart failure with reduced ejection fraction (HFrEF) or HFpEF [6, 7]. There are three different devices available for patients with either HFrEF or HFpEF: interatrial shunt device (IASD, Corvia Medical Inc., Tewksbury, MA, USA), V-Wave shunt (V-Wave Ltd., Caesarea, Israel) and Atrial Flow Regulator (AFR, Occlutech, Helsingborg, Sweden).

Atrial Flow Regulator is a self-expandable double-disc nitinol wire mesh construction allowing communication across the interatrial septum (Fig. 1). In contrast to V-wave and IASD, it is available in different sizes. The offered fenestration diameter ranges from 4 to 10 mm, but for HF patients only 8 mm and 10 mm have the European Conformité Européenne (CE) mark. Additionally, there are two available heights of the device: 5 and 10 mm, chosen according to interatrial septal thickness. The device is repositionable and retrievable.

First AFR implantation in Poland was done in a 28-year-old patient with severe pulmonary arterial hypertension (PAH) [8]. More recently, AFR has been successfully used as a bridge to lung transplantation in a young patient with drug-resistant idiopathic PAH [9].

Recently four AFR devices in patients with severe HFrEF were successfully implanted. These procedures were done as a part of ongoing PROLONGER trial (Pomeranian atRial flOw
Herein presented, are the results of a 66-year-old male with HFrEF (LVEF 25%), history of three myocardial infarctions, arterial hypertension and paroxysmal atrial fibrillation. Despite an optimal therapy, the patient remained symptomatic with New York Heart Association (NYHA) III. His 6-minute walk test distance (6MWT) was 200 m. The patient underwent diagnostic right heart catheterization with the Swan-Ganz catheter, which revealed decreased cardiac output accompanied with significantly increased pulmonary artery wedge pressure (PAWP).

According to PROLONGER protocol, hemodynamic indications for AFR are: PAWP above 15 mmHg or 25 mmHg at rest and exertion respectively. A right atrial pressure above 20 mmHg or exceeding PAWP is the contraindication for an atrial shunting procedure.

The AFR procedure was performed under general anesthesia. A three-dimensional transesophageal echocardiography (TEE) guided trans-septal puncture was performed followed by a 12 mm balloon septostomy. An AFR (8 mm fenestration, 5 mm height) was successfully implanted using a 12 F dedicated delivery system. The left to right mean gradient obtained from TEE was 2.7 mmHg. The patient was discharged home on the third day. As the patient had a history of atrial fibrillation, he was given non-vitamin K antagonist oral anticoagulants. There was no other specific indication for anticoagulation in this patient, because flow through the device was left to right.

The first follow-up visit after 1 month was complete, significant clinical improvement was noticed. The patient moved from NYHA III to NYHA II and 6MWT distance increased from 200 m to 397 m. Diagnostic right heart catheterization revealed significant reduction in PAWP, mean pulmonary artery and right atrial pressures (Fig. 2). The mean left to right gradient in TEE was 10 mmHg compared to 2.7 mmHg directly after AFR implantation. This difference could be explained by the fact that left atrial pressure had changed dynamically according to fluid overload, exertion and other conditions. Similar variability in all 4 patients thus far were observed.

The second follow-up visit was scheduled at 2 months.

This experience with AFR device is a promising option for patients with severe HF and further results will be published soon.

Conflict of interest: None declared

References


