Midterm results of sildenafil therapy in two complex patients with elevated pulmonary artery pressure after cavopulmonary connection

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Abstract

Two cases are presented. In the first patient (8-year-old boy) after Glenn operation without closure of pulmonary outflow from the common ventricle (SV-PA) despite transcatheter SV-PA closure increased mean pulmonary artery pressure (mPAP) (22 mm Hg) remained. After 6 months of sildenafil therapy he was catheterized again. His mPAP dropped to 10 mm Hg and a Fontan operation was subsequently performed. The second patient was a 25-year-old man, 20 years after a Fontan operation, presenting protein losing enteropathy and severe heart failure. All syndromes diminished significantly after medical therapy with sildenafil. Good clinical results of sildenafil therapy were maintained after 1 and 4 years of treatment. Chronic therapy with sildenafil can be beneficial in patients before and after the Fontan operation with elevated pulmonary artery pressure.

Key words: sildenafil, Fontan operation.

Introduction

Optimal therapy for patients with a functionally single ventricle still remains controversial. Surgical options to improve saturation of these patients are cavopulmonary connections (Fontan type operations). The main hemodynamic condition in such patients should be good ventricular function, low pulmonary pressure and low pulmonary resistance. The aim of this communication is to present midterm results of therapy with a pulmonary vasodilator – sildenafil in two patients with a single right ventricle after the Fontan or hemi-Fontan operation with elevated pulmonary artery pressure. Recently we discussed this problem in an internet edited issue of the medical journal Congenital Cardiology Today [1] and preliminary results of this therapy were published in Kardiologia Polska [2, 3]. Now mid-term follow-up of these patients is available.

Case reports

Case 1

A 7-year-old boy with mitral atresia, single ventricle, and malposition of great arteries was treated from the begin-
also treated with captopril, spironolactone and carvedilol. After 6 months he was catheterized again. His pulmonary pressure dropped more than half (mPAP 10 mm Hg), QP/QS was 0.43, PVR 2 Wood units. After 3 months he underwent a total cavopulmonary connection (TCPC) operation performed with an extracardiac tunnel (18 mm). The postoperative period was complicated with low cardiac output syndrome and right pleurothorax which needed 2 pleurocenteses. During the next days his clinical condition improved and he was discharged home 17 days after the operation on sildenafil (2 × 12.5 mg) and standard medication (captopril, furosemide, spironolactone, acenocoumarol). He remains in good condition after 1 year of follow-up [3].

Case 2

A 21-year-old male patient from another center was admitted to our department because of severe heart failure and protein losing enteropathy (PLE) as the consequence of failing Fontan. The initial diagnosis was mitral atresia, single ventricle and malposition of the great arteries. In infancy pulmonary artery banding and surgical atrial septectomy (Blalock-Hanlon) was performed. The second operation was performed at the age of 4 years (hemif-Fontan) and 1 year later, completion of Fontan – TCPC, without fenestration. During later follow-up he developed PLE with ascites. At the age of 15 years he was catheterized and mean pulmonary artery pressure (mPAP) was 27 mm Hg. Pulmonary artery anatomy was good. During the next 5 years his condition deteriorated – increasing peripheral swellings, ascites and cachexia. Because of ascites abdominal hernia developed which was closed surgically 1 month before his first admission to our department. At that moment there was still severe ascites with spontaneous drainage of peritoneal fluid from the postsurgical scar of the previously closed hernia. The circumference of the abdomen at that time was 130 cm with body weight 65 kg. He also had a visible dilated jugular vein, hepatomegaly (6 cm) and peripheral edema, desaturation (85%) and pleurothorax of the right lung. He was in NYHA class III. After examinations (ECHO, TC, MRI) good single ventricle function (EF 55%) and pulmonary anatomy was confirmed. During pleurocentesis 1400 ml of liquid was removed. Until that time he was treated with furosemide, hydrochlorothiazide, spironolactone, carvedilol and inibace. Because of no clinical improvement we decided to introduce sildenafil 3 × 25 mg daily. After 4 weeks of such therapy we observed spectacular clinical improvement (NYHA class II). He lost 5 kg of fluids, circumference of the abdomen was smaller (85 cm), the postsurgical wound closed successfully, and the liver diminished (3 cm). He was discharged home on this therapy. After 3 months he lost another 14 kg of fluid and his abdomen progressed to be smaller. Oxygen consumption test improved from 14 ml/kg/min initially to 28 ml/kg/min 1 year after sildenafil therapy. This clinical improvement persisted after 4 years of follow-up with only mild ascites and pleurothorax. Now he can continue his university studies. He refused control cardiac catheterization for assessment of pulmonary pressure [2].

Discussion

The administration of pulmonary vasodilators such as sildenafil has been shown to reduce elevated pulmonary artery pressure [4]. There is increasing evidence that at least half of post-Fontan patients have increased pulmonary vascular resistance as a consequence of pulmonary endothelial dysfunction. Goldberg et al. [5] suggested that maneuvers which increase cardiac output and lower central venous pressure can improve Fontan circulation. In case of PLE, with serum albumin level < 2.0 g/dl, he recommended treatment with sildenafil, CD-budesonite or Fontan surgical revision or heart transplantation. Moreover, the same author in a recently published paper showed that sildenafil may be a useful therapy to improve or maintain ventricular performance in selected patients after the Fontan operation [6]. Our observations in patients presented here as well as that of Deal and Jacobs [7] suggested that chronic pulmonary vasodilator therapy, in addition to chronic diuretics, may become part of routine long-term therapy in selected Fontan patients. What we noticed in our patients was not better ventricular performance in relation to sildenafil therapy but reduced pulmonary artery resistance resulting in clinical improvement. Meadows and Jenkins [8] in their comprehensive review summarized experience with evaluation, management and treatment of PLE in 18 patients from Boston Children Hospital (without application of sildenafil). Our cases and experience of others [9] indicate that sildenafil can be used safely and effectively in the treatment of patients with failing Fontan circulations. Interestingly, Ovaert et al. [10] in her study failed to show significant improvement after 3 months of treatment with bosentan (another type of pulmonary vasodilator) in 10 patients with failing Fontan.

In summary, patients with a single ventricle and elevated pulmonary artery pressure, both before and after the Fontan operation (failing Fontan), can benefit in our opinion from chronic sildenafil treatment.

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