The mid-aortic syndrome is a clinical condition generated by segmental narrowing of the abdominal or descending thoracic aorta. It might be either a congenital abnormality in the development of aorta or one of several acquired conditions such as infection, obliterator panarteritis, neurofibromatosis, retroperitoneal fibrosis, fibromuscular dysplasia, mucopolysaccharosis, and Takayasu’s arteritis. We are presenting a case of 17-years old female with life-threatening hypertension during the onset of pregnancy and unrecognized before the gestation middle aortic syndrome. Because of rarity of this syndrome, only ridiculous evidence exists with regard to the treatment. Single cases of surgery and interventional treatment with stent-graft placement have been reported, as well as good outcomes with only medical management. Unfortunately, the arterial hypertension in our patient was resistant on 3-drug antihypertensive regimen. She has successful endovascular mid-aortic syndrome repaired with CP covered stent on the 24-th week of pregnancy. The blood pressure decreased from 220/110 mmHg to 140/90 mmHg after the procedure. 3 months later the patient experienced the spontaneous vaginal delivery and gave birth to a healthy male newborn.

**Key words:** mid-aortic syndrome, pregnancy, endovascular stent graft placement.

**Introduction**

Coarctation of the abdominal aorta (CoA), or mid-aortic syndrome, is a rare vascular abnormality often producing life threatening arterial hypertension. Coarctation of the aorta accounts for approximately 11% of cases seen in the GUCH (grown-up with congenital heart disease) clinics and if not timely relieved is responsible for the persistent hypertension (30% at the age of 10 years; 66% at 40 years), premature coronary artery disease (25%) and severe berry aneurysms (2–3%). This arterial hypertension is usually resistant to all groups of the antihypertensive medications. Also, the hypertension is a frequent medical complication during pregnancy (normal pregnancy – preclampsia 11% in population). Unrecognized causes of the pre-existing hypertension occasionally become apparent only in pregnancy. Management of hypertension during pregnancy is challenging due to the fetal toxic impact of some drugs, namely ACE inhibitors and ABRs. Moreover, the females with aortic coarctation contemplating pregnancy should have repair prior to pregnancy. Management of hypertension in the uncorrected pregnant patient may be problematic, because of too low blood pressure below the coarctation site may result in compromising of fetus, aternal fetal death or abortion. The risk of aortic dissection or aneurysm rupture during pregnancy is low, but the maternal mortality risks are significantly higher if one of these occurs. To our best knowledge no one case of middle aortic syndrome repair during pregnancy has been described till nowadays. According to ESC guidelines on the management of cardiovascular diseases during pregnancy the percutaneous intervention for undiagnosed coarctation or re-CoA is possible during pregnancy, but it is associated with a higher risk of aortic dissection than in the non-pregnant patient and should only be performed if severe hypertension persists despite maximal medical therapy and there is the evidence of maternal or fetal compromise [3]. We are presenting a case of the endovascular treatment of mid-aortic syndrome with a CP covered stent during pregnancy, because of the uncontrollable hypertension.

**Case presentation**

A 17-years old woman admitted in her 24-th week of gestation with uncontrollable hypertension. On the physical examination the blood pressure was 220/110 mmHg on her arms. The blood pressure was resistant to three drug antihypertensive regimen: calcium channel blockers, β-blockers and thiazide diuretics. Also she had weak pulse on her legs. The lower limb pressure was unrecordable by the cuff measurements. The rest of the physical examinations were unremarkable. Coarctation of aorta was suspected.

Echocardiography showed mild left ventricular hypertrophy, the abdominal aortic flow was compatible with coarctation. During precise ECHO examination the coarctation of descending thoracic aorta above the diaphragm level was found with the peak gradient on it 90 mmHg (Figure 1). Because of pregnancy the patient was referred to MRI in order to get a better image and avoided the x-ray exposure. We used nonenhanced sequences with ECG-gated partial Fourier fast spin-echo, balanced steady-state free precession (SSFP). Noncontrast SSFP imaging enables rapid visualization of narrowing in «single shot» mode and a more detailed evaluation in any plane. The high signal-to-noise and contrast-to-noise ratio (due to cardiac and respiratory gating) and very quick sequences render SSFP useful for the pregnant patients. This investigation showed long segmental narrowing of descending thoracic aorta right above the diaphragm level with a pinhole orifice of about 2 mm in diameter (Figure 2).

It was decided to perform the percutaneous intervention for this patient because of the uncontrollable hypertension and major risk of the aortic dissection and possible rupture of the cerebral aneurysm. In the presence of a multidisciplinary team of an interventional cardiologist, cardiovascular surgeon, and obstetrician, specialized for cardio-vascular problems in pregnancy, neonatologist and anesthesiologist.

The procedure was performed under the local anesthesia. The right femoral artery was punctured. During all procedure the fetus was protected with a lead cover to negate the radiation exposure. Heparin was administered intravenously, at a dose of 100 IU/kg. 6 Fr multi-purpose catheter (Cordis, Johnson&Johnson, USA) was used to cross the stenosis in the aorta with the help of 0.035” glide wire (Terumo Company, Japan), which was advanced in a stable position across the stenosis and located in the ascending aorta. The hemodynamic measurements were recorded with the aortic pressures above and below the narrowing. The obtained gradient was 110 mmHg. The concentric narrowing of thoracic aorta above the diaphragm level was shown by the biplane aortography in the antero-posterior and lateral view (Figure 3).
There was also no evidence of the abdominal aortic branches stenosis due to this investigation. Measurements of the anatomic details were performed. The diameter of the aorta above and below the stenotic segment was 14 mm, the narrowest lesion was 2 mm. The stiff exchange length 260 mm 0.035” Amplatz guidewire (Cordis, Jonson&Jonson, USA) was positioned across the stenosis with a soft J-curve tip in the ascending aorta. The appropriate CP covered 8Z34 stent (NuMED Canada Inc.) was mounted on the 14x40 mm balloon-in-balloon (BiB) catheter (NuMED Canada Inc.) in a position related to the markers on the balloon catheters. 10 Fr Mullins sheath (Cook Medical, USA) with a radio-opaque marker at its tip was placed across the stenosis with the marker above the stenosis. The stent/balloon assembly was...
advanced through the sheath under fluoroscopic guidance and placed across the stenosis. The accurate positioning of the stent was checked with the hand injections of contrast through the side-port of the sheath. Inner and outer balloon inflation was subsequently performed up to 6 atm. without sufficient stent opening on the stenotic segment of the vessel. The Multitrack catheter (B. Braun Interventional Systems Inc., USA) was inserted over the exchange wire into the aorta above the dilated area and aortography was repeated in the same projection as prior to the stent implantation. The first aortography showed no changes after the stent-graft placement (Figure 4). Also, there was almost no pressure gradient change across the stenosis. It was decided to use the high pressure balloons. We repeated angioplasty with OptaPro 9x20 mm (Cordis, Jonson&Jonson, USA) and Atlas 12x20 mm (Bard Peripheral Vascular Inc. USA) with the pressure of 10 and 16 atm. consequently, repeating the aortography after each attempt. The final aortography showed a satisfactory result on the angioplasty site (Figure 5). Despite the slight recoil of aorta the gradient across stenosis dropped to 25 mmHg. The systolic blood pressure decreased to 140/90 mmHg. The decision was made to stop the procedure in order to avoid the high risk of aortic wall injury. The post procedural ECHO showed normal abdominal aortic flow (Figure 6).

After the procedure the patient’s blood pressure never again increased above 140/90 mmHg. According to the guidelines on arterial hypertension management during pregnancy, the aggressive treatment is mandatory when the blood pressure is higher than 170/110 mmHg [4]. So our patient had no necessity of any antihypertensive treatment, but the attention was paid for the early detection of any symptoms of the superimposed preeclampsia. The spontaneous vaginal delivery under the epidural anesthesia occurred in 39 weeks of pregnancy, a healthy male newborn 2700 g, 50 sm, 7–8 Apgar score was born. There was no evidence of any cardiovascular complication during the delivery period. Active management of the III period of labour was implemented, the early postnatal period was physiological. The newborn and
mother were discharged on the 7th day of the postnatal period. There was no evidence of any congenital heart diseases in the newborn by means of EchoCG performed on the 4th day of life.

Discussion

The mid-aortic syndrome is a life threatening condition because of possible complications caused by the upper limb arterial hypertension. Moreover, the native severe coarctation is a condition in which pregnancy risk — WHO IV (World Health Organization), what means — pregnancy contraindicated [3]. There was just a few cases of the mid-aortic syndrome diagnosed during pregnancy reported [1,2,5,6]. The secondary hypertension during pregnancy is present in 5 to 10% of cases and must be considered in presence of drug resistance [5]. Treatment of the arterial hypertension must be avoided in the patient with coarctation of aorta to prevent the placental hypoperfusion [3]. All antihypertensive drugs are assumed to cross the placenta and reach the fetal circulation. However, none of antihypertensive agents in routine use have been documented to be teratogenic, but trypyrine ACE inhibitors and ARBs are fetotoxic. So, the mentioned medication should be avoided in the pregnant woman [4].

Depending on the experience of the units different treatment strategies are used in the management of mid-aortic syndrome. The surgery is widely accepted especially in older patients and in the complex mid-aortic syndrome associated with the renal and visceral arterial stenosis. The type of surgery may involve a thoracoabdominal to infrarenal aortic bypass with the renal artery re-implantation, splenorenal bypass, aortorenal bypass, and autotransplantation. Experience over the last decade has shown that this pathology can be treated with the percutaneous techniques, such as balloon angioplasty or stent implantation, depending on anatomy and age of the patient [1,8].

The largest study with experience of treatment of the mid-aortic syndrome over last 30 years belongs to Boston Children’s Hospital and Harvard Medical School [8]. This study involved 53 patients. 35 pts underwent the invasive management, of them 21 pts had the percutaneous interventions, 5 pts — surgical techniques, 9 pts — both. According to this study the percutaneous interventions were very successful in decreasing the gradient across stenosis. However, freedom from reintervention at 1 and 5 year was 58% and 33% respectively after the percutaneous techniques, and it was longer after the surgery — 83% and 72% respectively.

But, to our best knowledge no one case of the mid-aortic syndrome repair during pregnancy has been described. There are some sporadic cases of the mid-aortic syndrome diagnosed during pregnancy, which was successfully managed just with medication [1,2]. As was previously stated, the percutaneous intervention for undiagnosed coarctation or re-CoA is possible during pregnancy, but it is associated with a higher risk of aortic dissection than outside the pregnancy and should only be performed if severe hypertension persists despite maximal medical therapy and there is a maternal or fetal compromise [3].

The represented case gives us a clear message: the mid-aortic syndrome could be successfully and safely managed with the percutaneous stent-graft placement during pregnancy when the benefits exceed the potential risk.

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Средний аортальный синдром представляет собой клиническое состояние, возникающее вследствие сегментарного сужения брюшной или нижней грудной аорты. Это может быть как врожденной аномалией развития аорты, так и следствием одной из приобретенных заболеваний, таких как: инфекция, облитерирующий панартрит, нейрофиброматоз, зачеревший фиброз, фиброзно-мышечная дисплазия, мукополисахаридоз и синдром Такаясу. В статье представлен случай 17-летней пациентки с опасной для жизни артериальной гипертензией во время наступления беременности и нераспознанным до беременности средним аортальным синдромом. Из-за редкости данного синдрома, очень сложно определить и найти правильный подход к его лечению. В статье указаны отдельные случаи хирургии и интервенционной терапии с установкой стент-графта, а также отмечены положительные результаты безоперационного ведения болезни. К сожалению, артериальная гипертония у нашей пациентки была устойчива к трехпрепаратному антигипертензивному лечению. Однако у пациентки было отмечено значительное улучшение эндоваскулярного среднего аортального синдрома после установки искусственного имплантата СР на 24-й неделе беременности. После проведенной процедуры артериальное давление снизилось с 220/110 мм рт.ст. до 140/90 мм рт.ст. Через 3 месяца пациентка самостоятельно родила здорового мальчика.

Ключевые слова: синдром среднего аортального синдрома, беременность, установка эндоваскулярного стент-графта.

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Середній аортальний синдром являє собою клінічний стан, який виникає внаслідок сегментарного звуження черевної або низхідної грудної аорти. Це може бути як вродженою аномалією розвитку аорти, так і наслідком одного з набутих захворювань, таких як: інфекції, облитеруючий панартрит, нейрофіброматоз, зачеревий фіброз, фіброзно-м'язова дисплазія, мукополісахаридоз і синдром Такаясу. В статті зазначені випадки хірургії та інтервенційної терапії з установкою стент-графта, а також відмічені позитивні результати безоперационного ведення хвороби. Нажаль, артеріальна гіпертензія у нашої пацієнтки була стійкою до трьохпрепаратного антигіпертензивного лікування. Однак у пацієнтки було відмічене значне покращення ендоваскулярного середнього аортального синдрому після встановлення штучного імплантата СР на 24-му тижні вагітності. Після проведеної процедури артеріальний тиск знісся з 220/110 мм рт.ст. до 140/90 мм рт.ст. Через 3 місяці пацієнка самостійно народила здорового хлопчика.

Ключеві слова: синдром середнього аортального синдрому, вагітність, установка ендоваскулярного стент-графта.

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Статья поступила в редакцию 24.08.2016 г.