#0001
EVALUATION OF RADIATION DOSES FOR PEDIATRIC PATIENTS DURING INTERVENTIONAL CARDIOLOGY PROCEDURES AT HAMAD GENERAL HOSPITAL, STATE OF QATAR.
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The estimated risks associated with radiation exposure are higher in children compared to adults. The use of fluoroscopy in diagnostic and interventional cardiac catheterizations being done in children requires accurate determination of the associated effective dose. Diagnostic procedures such as right and left heart studies (R&L Heart) and interventional procedures like closure of patent ductus arteriosus (PDA) or atrial septal defect (ASD), pericardial tap and balloon angioplasty of pulmonary or aortic valves, branch pulmonary arteries or coarctation of the aorta are among the commonest procedures done for the pediatric age group undergoing cardiac catheterization.

In this study the results of an analysis of doses recorded for 203 cases over 2 years (2013 and 2014) carried out in pediatric patients. The data likely represent the largest set of radiation doses recorded in children undergoing cardiac catheterization. The maximum Kerma Aria Product (KAP) recorded for those patients were 9779 cGy.cm². The maximum Cumulative Dose at the Interventional Reference point (CD_IRP) was also evaluated and found to be approximately 999 mGy. Body weight (BW) and body surface area (BSA) were also considered.

Materials and Methods: Two X-ray biplane fluoroscopy were used in this study. Patient's age, weight, height, gender, and procedure type and fluoroscopy time. Kerma Aria Product (KAP) and Cumulative dose data were recorded for 198 patients. The average pediatric age, weight and height were 3.03 years, 13.8 kg and 88.4 cm respectively. Peak voltage was 60.8kVp – 80 kVp.

Conclusion: Evaluation of KAP and CD_IRP doses are important indicators for the pediatric dose management and it is recommended to include all those data in patient's records. Body weight is an important factor in determining the radiation dose for children undergoing cardiac catheterization. Using a newer technology and adopt different imaging protocols (reducing the P/s and F/s) would lower the radiation dose without compromising the image quality.

#0002
TWO CASES OF TURNER SYNDROME WITH HYPOPLASTIC LEFT HEART MANAGED WITH HYBRID PROCEDURE
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Background: The presence of low birth weight, extra cardiac anomalies or genetic syndromes has been associated with poor outcome with Norwood procedures for palliation of Hypoplastic left heart syndrome (HLHS). Children with Turner syndrome and HLHS have a very high operative mortality and post-operative complications from persistent pleural effusions. Thus, for these high-risk patients, a hybrid approach to the Norwood operation can be an alternative palliation strategy. There is limited literature about outcomes of hybrid procedures in patients with Turner syndrome.

Methods: We present two cases of Turner syndrome with HLHS, who successfully underwent hybrid procedures as initial palliation for HLHS who subsequently underwent surgical repair achieving two-ventricle circulation.

Results: Two patients with Turner syndrome underwent hybrid Norwood palliation at 9 and 14 days of age. Weight at the time of intervention was 2320 grams and 2300 grams and both had placement of an 15mm x 5mm and 17mm x 7mm pre-mounted stent expanded and the pulmonary arteries were banded down to 3.5 mm diameter. These procedures were performed in the catheterization laboratory. The CHSS risk score was +17.93 and 15.35 (positive number favors a Univentricular repair, with the magnitude of the difference expressed by the number). In both children
there was growth of the left heart with successful uncomplicated 2nd stage procedure at 6 months and 10 months of age. One child had residual pulmonary artery stenosis post 2nd stage operation that required balloon pulmonary angioplasty. At 6 months follow up, both cases were doing well clinically and had normal function in echocardiogram. Both had mild sub-aortic stenosis with no obstruction or regurgitation.

**Conclusion:** The 2 patients in this series had successful initial palliation with a hybrid approach to the Norwood operation with no significant procedural complications or pleural effusions. Both sustained adequate growth of the left heart that subsequently allowed biventricular repair. Hybrid approach to the Norwood procedure carries the advantage of avoiding cardiopulmonary bypass and early aortic arch reconstruction and should be considered for palliation of high-risk patients with HLHS, especially with Turner syndrome.

**#0003**
**INITIAL EXPERIENCE OF ATRIAL SEPTAL DEFECT CLOSURE USING THE NEW GENERATION CARDIA ULTRASEPT IITM DEVICE IN MEXICO.**

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We present the initial experience in Mexico of atrial septal defect closure using the new Cardia Ultrasert II™ device. We present a series of 5 patients with ASD previously selected as candidates with favourable anatomy (less than 38mm defect, rims greater than 5mm) to be subjected for closure of the defect through interventionism treated in the period April-August 2016. Prospective, observational, transverse and descriptive study. The group included 3 female patients (60%) with a mean age of 10 ±2.12 years. The haemodynamic and anatomical data were as follows: pulmonary artery systolic pressure 25.2 ±3.5 mmHg, pulmonary to systemic flow ratio 2.78 ±0.52, sepal defect diameter 17.78 ±6.18 mm, expanded defect diameter 20.7 ±6.56 mm. All septal occluder were delivered successfully. No residual shunt evidenced by angiography and intracardiac echocardiography. At follow-up to one month, all patients showed complete closure of the defect and continuous decreased of right ventricular diastolic diameter (38.6 ±2.33 mm (Z-score 2.97 ±0.22) vs 34.26 ±3.13 mm (Z-score 2.46 ±0.33)) p=0.78. No complications at follow-up have been reported. The new generation of the Cardia Ultrasert II™ device is a good alternative to percutaneously treat atrial septal defect.

**#0004**
**RECONSTRUCTIVE SURGERY OF HYPOPLASIA OF THE AORTIC ARCH**

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**Objective:** to evaluate the function of baroreceptors in patients after different types of surgical correction of hypoplastic aortic arch.

**Materials and Methods:** In this prospective cohort study evaluated the results of surgical treatment of 54 patients who underwent surgical treatment for aortic coarctation. The patients were divided into two groups according to the method of correction of the defect: reconstruction with the use of a modified reverse plastic of LPA (group I, n=27) and reconstruction using the “extended” anastomosis (group II, n=27 patients).

**Results:** the Postoperative period of observation was 25 (21-30) months. Spontaneous sensitivity of the baroreceptors differed between groups and was significantly higher in group II is 11.6 (10.5; 12.6) vs 9.1 (8.2;10.1) in group I, p -0.04. The velocity of pulse blood flow was also higher in group II 7.7 (5.8;9) (m/s) -1 compared to 6.5 (5,4;7,1) (m/s) -1 in group I and differed between groups P = 0.04.

**Conclusions:** Reduced sensitivity of baroreceptors in patients after a modified reverse plastic of the left subclavian artery may be regarded as the method of choice in patients with coarctation and hypoplasia of the arch as a method of reducing the frequency of arterial hypertension in the late postoperative period.

**#0005**
**RECANNULATION OF LEFT PULMONARY ARTERY WITH RADIO-FREQUENCY PERFORATION AND STENT ANGIOPLASTY AFTER FAILED HYBRID STENT ANGIOPLASTY**

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An infant with pulmonary atresia and ventricular septal defect (VSD), s/p right-sided modified Blalock-Taussig shunt as a neonate, underwent VSD closure and placement of a 13 mm homograft right ventricle-to-pulmonary artery (RV-PA) conduit with augmentation of proximal branch pulmonary arteries at 9 months of age. After hospital discharge, there were concerns about flow into the left pulmonary artery (LPA), and he was referred for catheterization at 11 months of age, where he was found to have discontinuity of the left pulmonary artery (LPA). Transcatheter attempts at entering the LPA were unsuccessful, although he required stent angioplasty of right pulmonary artery branches. He was taken to the operating room (non-hybrid suite), where, after blind dissection and progressive dilation with Hegar dilators up to 3.5 mm, the surgeon positioned, and we deployed, a 16 mm long EV3 Intrastent Max LD stent mounted on an 8x20 mm Z-Med II balloon. A poor-quality C-arm angiogram, without the ability to record, demonstrated contrast to the end of the stent without flow in the LPA, indicating that the stent was extravascular. The patient was taken to cardiac catheterization lab, where simultaneous angiograms in the LPA stent and left pulmonary vein wedge injection demonstrated overlapping of the stent and the LPA. The Baylis Radio-Frequency system was used to perforate from the blind end of stent into the LPA, allowing for passage of the microcatheter into the LPA, followed by angiography in the LPA to confirm position. We then deployed a Palmaz Blue 7x15 mm stent over a V-18 wire, overlapping with the existing stent, resulting in patency of the LPA from the RV-PA conduit, and no extravasation of contrast. Follow-up echocardiograms, and
cardiac catheterization nearly one year later demonstrated unobstructed flow from the RV-PA conduit to the LPA.

#0006
FIRST EXPERIENCE OF INTRODUCTION OF ENDOVASCULAR METHOD TREATMENT OF CONGENITAL HEART DISEASES AT INFANTS IN THE FIRST YEAR OF BIRTH AT PEDIATRIC CARDIAC CENTER OF THE MINISTRY OF HEALTH OF AZERBAIJAN

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Objective: comparison of quality and quantity of the operated patients with critical congenital heart diseases by endovascular treatment method at newly opened cardio-surgical center and already famous and one of the leading centers of the world.

Methods: Endovascular treatment method of critical congenital heart diseases at infants.

Results: 32 patients under 1 year out of 209 patients with critical congenital heart diseases have been operated by endovascular treatment method since the opening of the Pediatric Cardio-surgical Center of the Ministry of Health of Azerbaijan in 2011. They were patients with such diagnoses as: closed interartial septum – 11 patients, balloon dilatation of the pulmonary artery – 80 patients, closed arterial duct – 75 patients, balloon dilatation of the coarctation aorta – 9 patients, balloon dilatation of the aortic valve – 12 patients, as well as such kind of complex congenital heart diseases as coarctation duct + balloon dilatation of the pulmonary artery – 15 patients, balloon dilatation of the pulmonary artery + closed interartial septum – 10 patients, perforation of the pulmonary artery + Paltayif PBV, balloon atrioseptostomy – 5 patients. Thus, lethality was 2.3 %. 438 patients out of 600 with critical congenital heart diseases have been operated by endovascular treatment method within 2007 – 2016 at the National Institute of Cardiovascular Surgery named after N.M. Amosov. They were with diagnoses as AoS+ 63; CoAo – 74; Sp – 58; Paltayif PBV TOF – 12; Rashkind – 163; Stent PBA/MARCA – 17; closed arterial duct – 15. Positive result has been obtained at 95% cases and the patients were discharged in satisfactory condition. Thereby, mortality was 5%.

Conclusion: The World Pediatric cardio – surgery of critical congenital heart diseases is based on the endovascular treatment methods; 70% of critical congenital heart diseases require palliative or radical endovascular intervention; wider range of endovascular methods of treatment of critical congenital heart diseases is conducted in the leading cardio-surgical centers; endovascular treatment of congenital heart diseases can be adopted quickly and efficiently in new medical centers supplied with relevant equipment and training of specialists in the leading cardiac clinics of the world.

#0007
SUCCESFUL TRANSCATETHER AORTO-RIGHT ATRIAL FISTULA CLOSURE USING AMPLATZER VASCULAR PLUG II

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Aorto cameral fistula are rare, usually found in right cavities, aquired fistula are more common than congenital, there are few published cases. We report a 7 year old girl refered for cardiac murmur and history of frequent respiratory disease, ecocardiogram evidenced color flow, turbulent, tortous between aorta and right atrium, 6 mm diameter, systolic maximum gradient 92 mmhg, and mild right cavities dilation. Cardiac cath was performed: aortic root pig tail injection revealed tortous fistula originated in right coronary sinus next to right coronary ostium, 6 mm diameter, and ended in posterior right atrial wall. Normal coronarigraphy. A 4 fr jr catheter is advanced to the more distal segment of the fistula and trough double guire maneuver the delivery sistem was placed, a vascular plug ii 12 mm, was accurately positioned, and realeased with mild residual shunt trough the device, no complications occurred.

Discussion and Conclusions: Aortocameral fistula closure is recomend even in assymopatis patients, due to low rate procedure complications, the risk of ventricule volume overload, bacterial endocarditis, pulmonary vascular disease, aneurysm formation and spontaneous rupture due to constant permeability we report a succesful percutaneous aorta right atrial congenital fistula closure.

Key words: Aorto cameral • fistula • amplater vascular plug ii

#0008
RISK FACTORS FOR AN ELEVATED PRE-FONTAN VENTRICULAR END-DIASTOLIC PRESSURE IN PATIENTS WITH SINGLE VENTRICLE CONGENITAL HEART DISEASE

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Introduction: Systemic ventricular end-diastolic pressure (SVEDP) is an important determinant of pulmonary artery pressure in those with Fontan circulation. Predictors of an elevated SVEDP have been incompletely identified in this population.

Methods: All patients who underwent Fontan operation at our center between 1/2009 and 12/2013 were retrospectively identified. SVEDP at pre-Fontan catheterization and other relevant patient variables were extracted. Analysis was performed to identify variables associated with pre-Fontan SVEDP.

Results: We identified 61 patients with mean age at pre-Fontan catheterization of 3.9 ± 1.6 yrs and 36 (59%) had a systemic right ventricle (RV). Pre-Fontan SVEDP was positively associated with
systemic ventricular systolic pressure (beta=0.4, p=0.004), aortic systolic pressure (beta=0.3, p=0.007), aortic mean pressure (beta=0.3, p=0.02), and decreased ventricular shortening (p=0.03). Compared to those with pre-Fontan SVEDP ≤ 7 mmHg, patients with SVEDP > 7 mmHg had higher average ventricular systolic pressure (85.0 ± 7.5 mmHg vs. 78.7 ± 8.3 mmHg, p=0.003), higher average descending aorta mean pressure (62.4 ± 4.9 mmHg vs. 58.6 ± 8.1 mmHg, p=0.03), and a higher incidence of decreased ventricular shortening (36% vs. 15%, p=0.07). The pre-Fontan SVEDP was similar between those with systemic RV (7.3 ± 2.0 mmHg) and systemic left ventricle (LV) (7.2 ± 1.8 mmHg) (p=NS). For those with a systemic RV, the SVEDP decreased significantly from pre-Stage 2 to pre-Fontan measurements (8.7 ± 2.6 mmHg vs. 7.3 ± 2.0 mmHg, p=0.02), but not for those with a systemic LV (7.8 ± 2.0 mmHg vs. 7.2 ± 1.8 mmHg, p=0.3).

Conclusions: In patients undergoing Fontan operation, pre-Fontan SVEDP was associated with decreased ventricular shortening and markers of systemic afterload. Systemic blood pressure may be an important determinant of SVEDP in this population. SVEDP decreased significantly after Stage 2 for those with systemic RV; but not for those with systemic LV; the systemic RV may be particularly vulnerable to pre-Stage 2 volume loading and benefit more than the LV from unloading at the stage 2 operation.

#0009 PEROUS TRANSCATHETER CLOSURE OF PERIMEMBRANOUS VENTRICULAR SEPTAL DEFECTS IN ONE WORKING GROUP, LONGTERM FOLLOW UP

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Our goal in this work was to evaluate the safety and efficacy of percutaneous transcatheter closure of ventricular septal defects (VSD), mostly perimembranous types (VSDpm) and long-term results. The VSD is the most common congenital heart disease. Transcatheter percutaneous closure have been a novel technique. Material and Methods: Between December 2004 and December 2013, 300 patients with medical record of VSD were admitted to our study, previously admitted to the cath lab at our center for percutaneous treatment of their VSD with various types of devices. All patients were followed until December 2013, 1 to 109 months. VSD type treated: perimembranous (VSDpm) 93.85 % and muscular (VSDm) 6.14%. The VSD measures before the procedure by echocardiography or at cardiac catheterization were 2 - 18 mm. Successful implantation of the device was 91.4 % in all attempted cases. The type of device used was Amplatzer 73.30 % and the Nit Occlud Coil 26.69 %. Complications were mostly minor, major complications were 2.49% including the late follow-up. They were complete AV block in 2 cases, 0.99 %; 2 cases need late surgery in the follow up secondary to the VSD closure procedure, 0.99 % and 1 case that required removal of the device in surgery because of Hemolysis 0.5 %. Conclusions: Percutaneous closure of VSD in experienced hands can be performed safely and successfully with low morbidity and mortality. Long-term results are good; percutaneous closure of VSD is less invasive and could be taken as a reasonable proven alternative in the treatment of perimembranous ventricular septal defects as well.

#0010 USE OF SYMMETRICAL HYPERIOM PERIMEMBRANOUS VSD FOR PERVERVentricular CLOSURE OF MUSCULAR SEPTAL DEFECT

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The percutaneous closure of the muscular interventricular septal defects is used in children with lightweight and it's incrementing with more promising results every time. The devices used have been occluders designed for muscular septum, which possess a length of 7 mm. We described a case which is treated with this closure mode through a new device designed for Perimembranous Ventricular Septal Defect in a 3 month old infant with 4.3 kg of weight and history of intrauterine growth retardation whom was diagnosed with the presence of muscular interventricular communication with length of 6 mm which is associated with the presence of aortic coarctation. It was taken to the surgery room and under general anesthesia a correction of aortic coarctation with terminal technical term extended by left lateral thoracotomy, sternotomy sequentially and interventricular communication approach per ventricular puncture was performed and a septal occluder device perimembranous of 8 mm designed for interventricular communication was placed achieving a complete occlusion of the defect.

The device used constitutes a feasible alternative in this patients and we consider the smallest waist length an advantage. Offering a more apt configuration further adapting the diameters of the interventricular septum in this age, which exposes less of the device's material towards the ventricular cavity.

Key words: Ventricular Septal defect • Perventricular closure • Hyperiom

#0011 EXPERIENCE IN MANAGEMENT OF AORTIC COARCTATION DIAGNOSED DURING PREGNANCY

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Introduction: Native severe coarctation is a condition in which pregnancy is at risk – WHO IV, which means pregnancy is contraindicated. Diagnosis of aortic coarctation is quite poor in developing countries. Pregnancy is not rare in patients with coarctation and possibility of existence of this abnormality should be considered in every case of hypertension occurring during pregnancy, especially in cases of drug resistance. Treatment of the arterial hypertension is mandatory when the blood pressure is higher than 160/90 mmHg.
Materials and methods: We are presenting seven cases of secondary arterial hypertension management in pregnant women due to aortic coarctation. One of the women had mid-aortic syndrome without involvement of visceral vessels and another one was diagnosed with hypoplastic transverse arch after coarctation repair. Mean age of patients was 25.7 ± 5.28 years, mean body weight was 69.57 ± 9.74 kg. Mean term of gestation at the time of diagnosis was 23.28 ± 5.76 weeks. Mean systolic blood pressure on admission was 175.71 ± 32.58 [from 140 to 240] mmHg.

Results. All patients received antihypertensive drugs. Mean SBP on medication was 147.86 ± 29.70 [from 110 to 200] mmHg. Four patients whose SBP was higher than 160 mmHg had percutaneous intervention for their coarctation. Three of them had coarctation stenting. The woman with transverse aortic arch hypoplasia had arch stenting when she was at the 15-th week of pregnancy. She experienced spontaneous rupture of the fetal membrane at the day of intervention which was managed conservatively. Spontaneous uneventful vaginal delivery occurred in three women who had intervention before labor. The patient with transverse arch stenting had caesarian section done due to the residual arterial hypertension after procedure. Three patients had coarctation stenting after childbirth. All patients with native coarctation were managed with caesarian section and strict blood pressure control. Of them one woman experienced acute aortic dissection type A on the day of caesarian section. On the same day she had coarctation stenting and supracorony ascending aorta replacement. All pregnancies were completed successfully with healthy babies born in term. Mean SBP after intervention was 126.42 ± 10.69 mmHg. Mean pressure gradient decreased from 55.0 ± 20.81 to 13.71 ± 8.79 mmHg.

Conclusion. Stenting of coarctation during pregnancy seems to be safe and effective option. There is no sufficient evidence still to draw definite conclusions about the optimal time of interventions. But, in our opinion it should be done before the labor due to high risk of cardiovascular complications despite strict blood pressure control. Interventions before 24-th week of gestation should be avoided as well in order to prevent miscarriages. Further multicenter investigations are warranted.

Key words: coarctation of aorta • pregnancy • arterial hypertension.

#0012
A NOVEL USE OF THE AMPLATZER VASCULAR PLUG III IN PERCUTANEOUS CLOSURE OF VENTRICULAR SEPTAL DEFECTS
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Background: Percutaneous ventricular septal defect (VSD) closure was first reported by Lock in 1988. Since then, the procedure has undergone many modifications to the technique and devices to avoid complications, especially to the conduction system. The search for an ideal device for VSD closure that simplifies the procedure and minimizes complications is still ongoing. Here we report the first use of the Amplatzer Vascular Plug III in VSD closure.

Methods: Charts and baseline electrocardiograms (ECG) of patients who underwent VSD closure using AVP III were retrospectively reviewed. VSD dimensions and other relevant measurements were obtained from intra-operative trans-esophageal echocardiograms (TEE) and angiography. The patients’ first post-operative echocardiogram and ECG were reviewed as well as latest follow-up if present.

Results. 16 patients (9 males, 7 females) underwent successful closure of VSD using the AVP III (13 were peri-membranous (pm), 3 were muscular). Median age was 5.2 (1.6 to 16) years and median weight 14.5 (8.7-52.2) Kg. The VSD size was 5.2 (2.6-10) mm on the left ventricular side and 4.2 (2.7-3.3) mm on the right ventricular side. There were no major complications to any of the patients. Procedure and fluoroscopy times were 140.2 (80-200) and 30.4 (13-48) minutes respectively. Only one patient had trivial residual shunt on next day post-operative TTE. One patient developed mild tricuspid regurgitation (TR) post VSD closure and 2 patients had resolution of previously present TR. None of the patients developed new conduction system abnormalities. Follow up is available in 6 patients. None of the patients had increased TR nor have any developed new conduction system abnormalities.

Conclusion: The AVP III’s oblong shape can be a good match for select pm-VSDs and small muscular VSDs. The small profile of the device’s wings minimizes interference with the aortic or tricuspid valves and the small central pedicle also decreases the risk of conduction system complications. The device can be delivered through a soft guide catheter, which can ease the manipulation of the device into the VSD. Our initial experience with this device to close VSDs is promising but long-term follow up is required.

#0013
TRANSCATHETER CLOSURE OF VENTRICULAR SEPTAL DEFECT USING DIFFERENT AMPLATZER DEVICE OCCLUDERS: INITIAL EXPERIENCE OF SOUHAG UNIVERSITY HOSPITAL
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Objective: To assess the challenges, feasibility, and efficacy of transcatheter closure of ventricular septal defect (VSD); perimembranous or muscular using different Amplatzer device occluders in initial experience of Souhag University hospital.

Patients/Methods: between 2013 to 2016, 26 patients (14 male, 12 female) underwent percutaneous closure of VSD using different Amplatz device occluders. After obtaining the size of VSD from the ventriculogram and TTE or TEE, a device of 2 mm larger than the narrowest diameter was chosen. The device deployed either by creation of arteriovenous loop or by retrograde arterial approach. The procedure was done under guidance of TTE or TEE. Follow up evaluations were done 1 month, 6 months, 12 months and yearly after procedure with transthoracic echocardiography and 12 lead electrocardiography. A retrospective review of the treatment results and adverse events was performed.

Results: Successful device placement was achieved in 25/26 of patients (96.2%). Median defect diameter was 6.7 mm (range 6 to 11 mm). Median weight was 21Kg (range 12 to 50). Median age was 7 years (range
2 to 15). Of 26 VSDs; there were 6 midmuscular, one apical muscular and 19 PM VSDs. Muscular VSDs were closed by muscular occluders but PM VSDs were closed by 10ADO 1, 3 ADOII and 6 muscular occluders. Three patients underwent successfully combined transcatheter intervention; one patient underwent ASD closure with PM VSD closure, the second patient underwent PDA closure with PMVSD closure and the third patient underwent balloon pulmonary valvuloplasty with midmuscular VSD closure. The device embolized in one case to right pulmonary artery (6.3%). Retrieval attempt was unsuccessful. The VSD occlusion rate was 69% at completion of the procedure, rising up to 83% at discharge and 97% during follow-up. Small residual shunts were seen at completion of the procedure, but they disappeared during follow-up in all except one patient. The median follow-up period was 8 months (range 1 to 15 months). Complete atrioventricular block (CAVB), major complication or death was not observed in our study.

Conclusions: Percutaneous closure of muscular or PM VSDs with different Amplatzer device occluders is safe and feasible with good mid-term outcomes, but large numbers of patients are required.

#0014
INTRAVASCULAR ULTRASOUND IN PULMONARY VEIN STENOSIS INTERVENTIONS
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Background: Pulmonary vein stenosis (PVS) is an aggressive disease with high rates of morbidity and mortality. Surgical and catheter interventional approaches have yielded modest success, at best. Refinements in catheter intervention could potentially improve outcomes.

Methods: Single-center, retrospective review of patients with PVS undergoing cath from 3/2015 – 8/2016. As part of the diagnostic cath, the left atrium was entered via an existing septal defect or by transseptal puncture. Systemic heparinization was provided to maintain ACT>250. Intravascular ultrasound (IVUS) of the pulmonary veins was performed using an Eagle Eye® Platinum IVUS catheter (Volcano Corp, Rancho Cordova, CA).

Results: 3 patients underwent 4 catheterizations (1 diagnostic, 3 interventional). Median age was 2.2y (0.7-47.5y), weight 9.9kg (7.3-61kg). For the interventional caths, mean PV gradient was 10 mmHg with reduction by 8.8 mmHg. Two patients had congenital PVS, one was post-repair of Scimitar syndrome with an obstructed pulmonary venous baffle. For patient 1 (Fig. 1a), IVUS showed interval vessel growth (arrowheads) around a previously placed stent (arrow) allowing for redilation with good stent apposition after intervention (Fig 1b). On follow-up cath, stenosis was seen in areas no longer covered by stent material, due to foreshortening during prior redilation, which improved with angioplasty and additional stent placement. For patient 2, IVUS confirmed long-segment hypoplasia that was unlikely to respond to intervention. For patient 3, narrowing in the midportion of the obstructed Scimitar baffle was seen (Fig. 2a, arrow) with good stent apposition after intervention (Fig 2b). All patients recovered well from their procedures and there were no thrombotic complications from performing IVUS.

Conclusions: IVUS of the pulmonary veins is safe and easy to perform and provides very detailed imaging of PVS to help guide therapy. For those requiring intervention, adequate stent apposition to the pulmonary vein walls as well as limiting vessel overdilation may minimize future instent stenosis and need for reintervention in this challenging disease.
#0015

DEVICE TO THE RESCUE
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9 year old girl, presented with palpitation and progressive shortness of breath.
Underwent ICR for TOF 3 years back.
Echo showed a residual perimembranous VSD measuring 8 mm.

#0016

PERCUTANEOUS CLOSURE OF PATENT DUCTUS ARTERIOSUS – 20 YEARS EXPERIENCE OF ONE CENTER
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Introduction: Percutaneous closure has become a method of choice in treatment of patent ductus arteriosus (PDA). Recently different types of occluders have evolved. We present our own experience in this field.

Materials and Methods: Between 1996 and 2016 840 patients (pts; 543 f; 0.3-84.5y; median 4y) had percutaneous PDA closure performed in our center. PDA A type was present in 415 pts, B in 23 pts, C in 60 pts, D in 105 pts, E in 192 pts, G in 25 pts, moreover, 20 pts had previously ligated PDA recanalized. Reversible pulmonary hypertension (PH) was observed in 121 pts (14.4%). We divided this period in double umbrella (DU) era (1996-2000), detachable coils (C) era (1996-2014) and nitinol wire mesh devices era: like Amplatzer Duct Occluder type I (ADO I) (1997-now), Amplatzer Duct Occluder type II (ADO II) (2009-now) and Amplatzer Duct Occluder type II Additional Sizes (ADO II AS) (2014-now). Application techniques of mentioned above devices were routine. In special situations another devices were used.

Results: We have applied DU in 25 pts with 88% success rate (1 embolisation), C in 463 pts with 96.8% success rate (7 embolisations), in 250 pts ADO I and like ADO I occluders with 100% success rate (ADO I in 140 pts, Cera Occluder in 8 pts, Cardi-o-Fix in 64 pts, HeartR in 27 pts, Hyperion in 11 pts), ADO II in 15 pts with 100% success rate and ADO II AS in 71 pts with 98.6% success rate. Moreover, in type B of PDA 5 CardioSEAL/STARflex devices and 2 ASD nitinol wire mesh occluders; in type D of PDA 3 Amplatzer Vascular Plugs type II and in 6 pts with higher PH Muscular Amplatzer VSO were used with 100% success rate. Residual shunt 24 hours after the procedure was observed in 24% of DU, 16.7% of C and no residual shunts were observed in other groups of pts.

Conclusions: PDA percutaneous closure methods have quickly developed and nowadays they are not only safe and efficient but representing also relatively low complication rate.
#0017
TRANSCATHETER CLOSURE OF POSTINFARCTION VENTRICULAR SEPTAL DEFECT – IN-HOSPITAL OUTCOME AND FOLLOW-UP ANALYSIS
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Introduction: Postinfarction Ventricular Septal Defect (PIVSD) is a rare and severe complication with poor prognosis. Transcatheter closure (TC) of such defect can be a good alternative to surgery in selected patients.

Materials and Methods: All of 26 consecutive patients (pts) (64.6±10y; 9 female) in whom TC of PIVSD was attempted in our department between 2000-2015 were retrospectively analyzed. All pts had coronary artery angiography before TC of PIVSD. Mean time between PIVSD occurrence and its TC was 11±5 weeks. All pts were admitted in III/IV NYHA class, 9 pts were on balloon counterpulsation; 3 pts were after previous PIVSD cardiac surgery with residual shunt. Mean PIVSD diameter was 11.4±3.8mm (5-19mm) in angiography.

Results: 20 from 26 attempted PIVSD TCs were successful (74%) with immediate significant clinical improvement. In 6 pts procedure was abandoned because of unfavorable morphology, from whom 3 pts were referred for surgery. During the TC 4 pts needed defibrillation because of VF. No peri-procedural deaths were observed. Hemolysis occurred 2 days after TC in 1 pt. 15 from 20 pts survived till the discharge; 5 pts died because of multiorgan failure. In observation 2 pts occurred 2 days after TC in 1 pt. 15 from 20 pts survived till the discharge; 5 pts died because of multiorgan failure. In observation 2 pts needed second percutaneous closure of another defect. Mean follow-up was 6±4,6y (11 pts).

Conclusions: PIVSD is one of the most severe complication of myocardial infarction with high risk of surgical or medical treatment. TC of such defect should only be limited to properly selected pts.

#0018
COMPARISON STUDY OF THREE-DIMENSIONAL TRANSESOPHAGEAL ECHOCARDIOGRAPHY AND CT IN MEASURING THE SIZE OF AORTIC RING AND THE HEIGHT OF CORONARY
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To investigate the difference between three-dimensional transesophageal echocardiography (3DTEE) and CT in measuring the size of aortic ring and the height of coronary ostium.

Methods: 15 patients were recruited and were treated with the transcatheter aortic valve implantation (TAVI). Routine transthoracic echocardiography, two-dimensional and three-dimensional echocardiography (2DTEE, 3DTEE) and CT examinations were taken preoperatively.

Results: The minimal diameter, maximum diameter, perimeter, area of the aortic ring measured by 3DTEE showed strong correlation and consistency with those measured by CT [r=0.88, P<0.0001, ICC=0.928 (0.788-0.976); r=0.81, P=0.0003, ICC=0.890 (0.673-0.963), r=0.85, p=0.0001, ICC=0.914 (0.744-0.971); r=0.88, P<0.0001, ICC=0.932 (0.799-0.977)]. The height of the left and right coronary artery measured by 3DTT also showed strong correlation and consistency with those measured by MDCT [r=0.87, P<0.0001, ICC=0.923 (0.777-0.975); r=0.82, P<0.0002, ICC=0.897 (0.693-0.965)]. Besides, inter-observer and intra-observer reproducibility for 3DTEE measurement data were very good.

Conclusions: 3DTEE has high repeatability in evaluating minimal diameter, maximum diameter, perimeter, area of the aortic ring and the height of coronary ostium, which also shows good correlation with those measured by CT. Thus, 3DTEE is expected to replace CT in the near future.

#0019
STENT PLACEMENT FOR TREATMENT OF AORTIC COARCTATION IN CHILDREN UNDER 30 KG: ACUTE AND LONG-TERM OUTCOMES
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Introduction: The use of stents for percutaneous treatment of aortic coarctation (CoAo) in children presents some challenges as possible limitations of vascular access and lack of published data regarding the need for reintervention. We sought to evaluate the feasibility, safety and efficacy of stent in children, with an emphasis on follow-up.

Methods: A retrospective analysis of a cohort of consecutive children (<30 kg) who had undergone stent placement for aortic coarctation between April/2009 and December/2015 was performed. The access route was the carotid in patients <10 kg. Stents that can be expanded between April/2009 and December/2015 was performed. The access route was the carotid in patients <10 kg. Stents that can be expanded to larger diameters have been used. Demographic, clinical, hemodynamic and follow-up data were collected. The endpoints evaluated included: immediate and mid-term severe adverse events (SAE), persistent high blood pressure and need for reintervention.

Results: Thirty-seven patients (25 male, 27 with native CoAo and 1 with associated hypoplasia of aortic arch), with mean age and weight of 5.4±3.4 years and 20.7±11.0 kg were enrolled respectively. Immediately, the peak-peak gradient decreased from 33.7±15.1 to 5.4±5.3 (p <0.01) and the ratio of aortic coarctation diameter/DAo increased from 0.40±0.16 to 0.95±0.20 (p <0.01). There were no deaths or immediate SAE. Thirty-four patients were followed during a mean period of 43.1 ± 19.4 months. Computed tomography of aorta was performed in 26 patients (23.1 ± 17.6 months after procedure) and showed stent integrity in all. Five patients still needed medication

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for high blood pressure. Seven patients required percutaneous reintervention (36.1 ± 19.0 months after initial treatment) due to aortic aneurysm (1), residual stenosis above the stent (1) and adjustment to somatic growth (5). One patient required surgery due to residual hypoplasia of aortic arch (15.1 months later). All of reinterventions have been carried out successfully without any SAE.

**Conclusion:** Stenting for treatment of CoAo in children was feasible, safe and effective in reducing blood pressure levels and gradient. A significant rate of reintervention was observed because of the previously known need to stent adjustment to somatic growth in most cases (SAE were rare). On the same way, post dilation of stents for CoAo in children has proved to be feasible and effective.

**#0020**

**PARTICLE EMBOLIZATION OF SYSTEMIC-TO-PULMONARY COLLATERAL ARTERY NETWORKS IN CONGENITAL HEART DISEASE: TECHNIQUE AND CONSIDERATIONS**

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**Background:** Systemic-to-pulmonary artery collateral (SPC) networks commonly develop in patients with single ventricle physiology and chronic hypoxemia. Though these networks augment pulmonary blood flow, much of the flow is ineffective which contributes to cardiac volume loading. This volume loading can have detrimental effects, especially for single ventricle patients. Some data suggest that occluding these collaterals may improve outcomes after subsequent operations, especially when the volume of collateral flow is significant. For other patients—e.g. with hemoptysis—collateral occlusion is a crucial therapy. Traditional practice has been to coil occlude the feeding vessel, but this technique has limitations.

**Cases & Technique:** We reviewed all procedures that included SPC embolization from August 2013 to June 2016. We perform particle embolization utilizing a co-axial catheter system to deliver 510-700 micron particles deep into feeding arteries (figures).

**Results:** We performed particle embolization during 42 catheterizations on 34 patients. The majority of patients had single ventricle physiology; a few patients presented with hemoptysis or had incidentally noted SPCs. Particle embolization was acutely successful with the majority (93%) having no residual flow. No complications occurred.

**Discussion:** Traditional coil occlusion of SPCs is suboptimal for multiple reasons. First, occlusion of fistulous connections is ideally performed as distally as possible. Coil occlusion generally occludes the most proximal source. Second, SPCs tend to recur from the same feeding vessel. Coil occlusion prevents re-access of the feeder, impeding occlusion of recurrent SPCs. Particle occlusion avoids both these limitations as the technique occludes distal connections while maintaining patency of the feeder vessel. The occlusion is therefore more immediately effective—by occluding distal connections—and avoids problems with re-accessing feeders. Thus, embolization with particles may be a superior technique than coil occluding the proximal feeding vessel.

**#0021**

**RETRIEVAL OF A FIGULLA OCCLUTECH SEPTAL OCCLUDER EMBOLIZED DEVICE FROM RIGHT VENTRICLE USING ITS NATIVE DELIVERY SYSTEM. CASE REPORT OF A NOVEL APPROACH**

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A 27-year-old lady was diagnosed as having a large secundum atrial septal defect and moderate pulmonary artery hypertension on echocardiogram. She was planned for transcatheter device closure of atrial septal defect as transesophageal echocardiography revealed ASD secundum (31 mm × 29 mm) with adequate rims. The patient was taken to the catheterization laboratory for transcatheter closure under local anaesthesia. A 14 F Cook sheath was selected to deliver 33 mm device. The device was loaded on to the delivery system and delivered across the defect using the right upper pulmonary vein technique. The device fitted nicely on to the septum and final position confirmed on TOE and released. The next morning to confirm position of device on TTE before discharging the patient, it was revealed that the device had embolized in to RV. The patient was again taken to the cath lab and after multiple attempts with a 15 mm Amplatzer gooseneck snare (ev3 Endovascular Inc., Plymouth, MN, USA) the device could not be pulled in to the sheath satisfactorily. Finally we used native delivery cable within 14 F sheath. After few attempts we were able to retrieve the 33 mm ASD device with locking mechanism of delivery cable. Then we closed the same ASD with 36 mm occlutech septal occluder with no further complication. The patient was discharged next day with advise of tab loprin 150 mg daily for 6 months.

**Conclusion:** In certain situations of embolized occlutech devices, native delivery cable with two prongs can be very effective and safe with stability to retrieve when snare is unable to capture and stabilize. However the capturing area of prongs is much less than snare.

**#0022**

**SUCCESSFUL HEMOSTASIS OF ACUTE LUNG BLEEDING USING AMPLATZER VASCULAR PLUG AND COILS IN A PATIENT WITH PULMONARY HYPERTENSION**

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Lung bleeding is the dreadful complication of cardiac catheterization that can directly result in demise in patients with pulmonary hypertension. We report successful hemostasis achieved by combined occlusion of Amplatzer Vascular Occluder and coils.

**Case:** A 61-year-old female patient who is known to have pulmonary hypertension with mean pulmonary artery pressure of 66 mmHg underwent follow-up diagnostic catheterization. She had a large patient arterial duct with a diameter of 12 mm treated with Amplatzer Septal Occluder 7 months ago and her past history includes pulmonary tuberculosis resulted in chronic respiratory insufficiency that required home oxygen treatment.
When we tried to measure right pulmonary wedge pressure, she suddenly complained chest pain and showed massive hemoptysis that progressively suffocated her. She fell in shock, but was successfully treated by volume expansion, intravenous adrenalin infusion, and placement of laryngeal mask successfully replaced with blind tracheal intubation.

Repeated exploratory pulmonary angiography showed unconfined pulmonary bleeding that could lead to continuous bleeding without treatment. We placed a 4 French Judkins left coronary type catheter into right lower branch and placed 2 Flipper coils with loop-diameters of 6.5 mm and 5 mm. However, there was still continuous bleeding and we place a Amplatzer Vascular Plug of 8 mm in diameter that could successfully brought complete occlusion and stop pulmonary bleeding.

Conclusion: Cardiac catheterization of patients with pulmonary hypertension carries a risk of severe complication, namely pulmonary bleeding. Preparation of laryngeal mask and occlusion devices such as coils and Amplatzer vascular plugs are mandatory in this setting.

#0023 INTRA-CARDIAC ECHOCARDIOGRAPHY GUIDED TRANS-CATHETER CLOSURE OF PATENT DUCTUS ARTERIOSUS WITHOUT CONTRAST ANGIOGRAPHY
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Background: Though contrast angiography is the standard guidance of trans-catheter closure of patent ductus arteriosus (TC-PDA), it is contra-indicated in patients with severe renal disease that often seen in senile patients. We have developed intra-cardiac echocardiography (ICE) guided TC-PDA (Cathet Cardiovasc Intervent 2015). We report sequential 6 cases that successfully underwent TC-PDA without contrast angiography.

Materials and Methods: Subjects were 5 patients with PDA and median age of 57.4 (35.4 - 66.1) years old. The median size of PDA was 4.6 (3.2 - 11.7) mm with median Qp/Qs of 1.9 (1.4 and 2.4), respectively. The oldest patient suffered from renal dysfunction and 2 patients had pulmonary hypertension. Prior to the TC-PDA, all patients underwent contrast X-ray computed tomography to clarify the anatomy. ICE catheter was inserted through 2nd sheath at femoral vein and placed at main or left pulmonary artery. During the TC-PDA, we primarily used ICE to guide the procedure.

Results: We could successfully place Amplatzer Duct Occluders in 4 and Amplatzer Septal Occluder in 1 without contrast angiography. ICE at main or left pulmonary artery has allowed us to determine the diameter and length of PDA, to monitor the device placement, and to determine the residual shunts. ICE did not increase the risk of complication except for transient arrhythmia, though new operator needs some learning time to understand orientation of ICE.

Conclusion: ICE-guided TC-PDA without contrast angiography is feasible and can be the standard treatment for adult patients with renal dysfunction.

#0024 COMPLICATIONS FOLLOWING PERVERVENTRICULAR DEVICE CLOSURE OF MUSCULAR VENTRICULAR SEPTAL DEFECT
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A male infant was born with tricuspid valve (TV) dysplasia with severe regurgitation and Swiss-cheese muscular ventricular septal defects (MVSD). Diagnostic cardiac catheterization demonstrated significant left-to-right shunting with a Qp:Qs ratio of 2.29:1. He underwent attempted perventricular device closure of the MVSDs with surgical repair of the TV. A purse-string was placed into the mid-portion of the right ventricle (RV) free wall to provide direct access to the anterior MVSD. Attempts to close the apical-MVSD with a device were unsuccessful due to its crowding with RV trabeculations. Hence, the decision was made to leave the apical-MVSD without further intervention. Transesophageal echocardiogram suggested a small outpouching posterior to the left ventricle (LV) free wall measuring about 5x7mm that was concerning for a potential LV-PSA (pseudaneurysm) caused by the wire vs. sheath across the VSD. Therefore, cardiopulmonary bypass was initiated, the TV was repaired and a MPA band was placed.

An echocardiogram showed enlargement of the LV-PSA to 3.7x4.0x2.7 cm, which was confirmed with cardiovascular magnetic resonance, revealing a narrow neck at its origin from the LV apex. Due to the concern for rupture, the aneurysm was repaired surgically. The two devices were extracted and the aneurysmal sack was completely resected.

#0025 INITIAL EXPERIENCE OF ATRIAL SEPTAL DEFECT CLOSURE USING THE NEW GENERATION CARDIA ULTRASEPT IITM DEVICE IN MEXICO.
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We present the initial experience in Mexico of atrial septal defect closure using the new Cardia Ultraspe II™ device. We present a series of 9 patients with ASD previously selected as candidates with favourable anatomy (less than 38mm defect, rims greater than 5mm) to be subjected for closure of the defect through interventionism treated in the period April-August 2016. Preliminary prospective, observational, transverse and descriptive study. The group included 7 female patients (76%) with a median age of 8 years (1-13). The hemodynamic and anatomical data were as follows: pulmonary artery systolic pressure 25.14 ±3.9 mmHg, pulmonary to systemic flow ratio 2.38 ±0.66, septal defect diameter 17.78 ±6.18 mm, expanded defect diameter 22.6 ±5.82 mm. All septal occluder were delivered successfully. No residual shunt evidenced by angiography and intracardiac echocardiography. At follow-up to 2.1 months, all patients showed complete closure of the defect and continuous decreased of right ventricular diastolic diameter with an initial median of 38mm (30-40) and after catheterization of 28.5mm (23-30), p=0.01 and Z-score of 3 (2.87–3.15) vs 1.8 (1.5–1.95), respectively, p=0.01. The new generation of the Cardia Ultraspe II™ device is a good alternative to percutaneously treat atrial septal defect.
Veno-venous collaterals (VVC) may lead to significant systemic desaturation after Fontan operation. We present a 16-year-old girl with univentricular physiology who had undergone palliation with a Damus-Kaye-Stansel procedure, Glenn operation and Fontan completion at another institution. Cardiac catheterization had revealed mean Fontan pressures of 23 mmHg. Subsequent treatment with Bosentan had resulted in a drop to 15 mmHg several years before. A large VVC from the innominate vein to the left atrium had been identified with systemic SaO2 ranging from 85-90%. Interventional closure had been attempted several times at three different large volume pediatric cardiac centers. Finally, this patient was transferred to our institution in severe heart failure with ascites, systemic desaturation and poor ventricular function. Under mechanical ventilation with FiO2 1.0 and NO 20 ppm, arterial SaO2 could not be increased above 60%. Despite optimal medical management, her status deteriorated rapidly. A single chamber cardiac assist device (Berlin Heart) was implanted and the girl was listed for urgent heart transplantation. Due to continued severe hypoxemia and metabolic acidosis it was decided to close the large VVC via a hybrid approach as other access routes had failed before. Access via the left ventricle was chosen in order to achieve a straight angle to the insertion of the VVC into the roof of the LA. After introduction of a sheath into the LV, the VVC could be entered with a right Judkins catheter and an 8F Amplatzer sheath could be advanced deep into the VVC subsequently. On angiography the VVC measured 12 x 14 mm in diameter at its narrowest portion, the widest site was 22 mm. Accordingly, an 18 mm Amplatzer Vascular Plug (AVP) II was deployed in the distal VVC. For complete closure a second AVP II (20 mm) was implanted slightly more proximally. Arterial oxygen saturation immediately increased to 80% thereafter. Final angiogram demonstrated adequate and stable closure at another institution. Cardiac catherterization had revealed mean Fontan pressures of 23 mmHg. Subsequent treatment with Bosentan had resulted in a drop to 15 mmHg several years before. A large VVC from the innominate vein to the left atrium had been identified with systemic SaO2 ranging from 85-90%. Interventional closure had been attempted several times at three different large volume pediatric cardiac centers. Finally, this patient was transferred to our institution in severe heart failure with ascites, systemic desaturation and poor ventricular function. Under mechanical ventilation with FiO2 1.0 and NO 20 ppm, arterial SaO2 could not be increased above 60%. Despite optimal medical management, her status deteriorated rapidly. A single chamber cardiac assist device (Berlin Heart) was implanted and the girl was listed for urgent heart transplantation. Due to continued severe hypoxemia and metabolic acidosis it was decided to close the large VVC via a hybrid approach as other access routes had failed before. Access via the left ventricle was chosen in order to achieve a straight angle to the insertion of the VVC into the roof of the LA. After introduction of a sheath into the LV, the VVC could be entered with a right Judkins catheter and an 8F Amplatzer sheath could be advanced deep into the VVC subsequently. On angiography the VVC measured 12 x 14 mm in diameter at its narrowest portion, the widest site was 22 mm. Accordingly, an 18 mm Amplatzer Vascular Plug (AVP) II was deployed in the distal VVC. For complete closure a second AVP II (20 mm) was implanted slightly more proximally. Arterial oxygen saturation immediately increased to 80% thereafter. Final angiogram demonstrated adequate and stable closure.

The data obtained demonstrates how relatively small changes in our practice and radiation awareness can decrease the amount of radiation exposure to the patients and staff in the cardiac cath lab. Our program will continue to remain aware of radiation exposure and strive to keep the dose as low as possible. With the guidance of the National Cardiovascular Data Registry (NCDR®) Impact Registry™, we are able to maintain and keep track of our improvements.
requiring anesthetic medication boluses) and lack of failed sedation (need to convert from CS to GA). Multivariate logistic regression models were utilized to compare results.

Results: Of 104 infants who underwent a pre-stage II catheterization, 82 (79%) had safe sedation and 56 (54%) had effective sedation. CS was utilized in 91 (88%) patients and 8 (10%) required conversion to GA (failed sedation). There were no differences between CS and GA patients in baseline demographics, shunt type, procedural duration, intra-procedural lowest pH/highest PCO2, or rates of safe and effective anesthetic management. However, ICU admission was more common in GA patients (23% vs 2%, p=0.013) and patients with higher intra-procedural PCO2 (OR 1.17, p=0.013). Higher PCO2 was also associated with greater odds of a failed sedation (OR 1.18, p=0.004). Higher baseline oxygen saturation was independently associated with a safe catheterization (AOR 1.12, p=0.014) and higher weight was independently associated with both safe (AOR 2.86, p=0.004) and effective (AOR 1.74, p=0.013) anesthetic management.

Conclusions: CS can provide safe anesthetic management for SV infants undergoing a pre-stage II diagnostic catheterization and few patients require conversion to GA. However, hemodynamic steady state can be difficult to achieve regardless of the anesthetic management strategy.

#0029
FIRST HUMAN CASES OF SELF-EXPANDABLE PERCUTANEOUS PULMONARY VALVE IMPLANTATION USING KNITTED NITINOL-WIRE STENT MOUNTED WITH A TRI-LEAFLET PORCINE PERICARDIAL VALVE
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Background: Severe pulmonary regurgitation (PR) and associated right ventricular (RV) dilatation in native right ventricular outflow tract (RVOT) is challenging and still on clinical trial. We report first human cases of self-expandable percutaneous pulmonary valve implantation (PPVI) using newly made knitted nitinol-wire stent mounted with a tri-leaflet porcine pericardial valve developed in South Korea.

Methods: We reviewed 8 cases of self-expandable PPVI at the Seoul National University Children's Hospital. This self-expandable valved-stent was newly developed by our research team with the cooperation of the TaeWoong medical company in South Korea. This valved-stent was made by knitted nitinol-wire backbone with tissue valve using porcine pericardium with multiple steps for tissue preservation including decellularization and alpha-galactosidase treatment.

Results: Eight patients underwent total correction of Tetralogy of Fallot previously and showed severe PR (mean PR fraction: 43.7%, range: 35.4-56) and enlarged RV volume (mean indexed RV end-diastolic volume: 188.6 mL/m², range: 167.5–209.8). Their median age at PPVI was 21 years old (range: 13-26). At the targeted RVOT area, 4 patients were implanted with 28 mm diameter valved-stent and 4 patients were implanted with 26 mm diameter valved-stent loaded in the 18 French delivery cable. There were no significant peri-procedural complications in all patients. After procedure, there was no significant pulmonary stenosis or PR from cine-angiography and echocardiography in all patients. Chest X-ray showed good valved-stent position at targeted RVOT area. All patients discharged 4 days after PPVI without any problem. One patient completed 6 months follow-up after PPVI and showed decreased indexed RV end-diastolic volume from 181.7 to 126.7 mL/m² from cardiac MRI.

Conclusion: First human implantation of self-expandable percutaneous pulmonary valve using knitted nitinol wire mounted with a tri-leaflet porcine pericardial valve developed in South Korea was feasible and effective at short-term follow-up. A clinical trial for feasibility to evaluate the safety and short-term effectiveness of this self-expandable valved-stent for 10 patients is ongoing for the congenital heart disease with pulmonary valve disease in South Korea.

#0030
SAFETY OF OUTPATIENT CARDIAC CATHETERIZATION IN INFANTS WITH SINGLE VENTRICLE CONGENITAL HEART DISEASE.
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Background: The benefits of outpatient cardiac catheterization (cath) were first assessed in the 1980s; it reduces anxiety to patients and families and decreases cost. Cardiac cath is routinely performed in patients with single ventricle congenital heart disease (SVCHD) to aid in hemodynamic assessment, intervention and surgical planning. There is significant morbidity and mortality associated with interstage SVCHD in shunt-dependent patients and substantial intra-procedural variation exists between centers. Post-procedural best practices following cardiac cath of infants with SVCHD are unknown. Our institutional strategy has been to discharge patients following a 4-6 hour post-procedural observation period. The objective of this study was to investigate the incidence and causes of readmission of infants with SVCHD following outpatient cardiac cath. Methods: We performed a retrospective review of all patients less than one year of age with SVCHD who underwent cardiac cath between 2007 and 2015 at our institution querying the Society of Thoracic Surgeons Database. Unplanned readmissions were defined as an admission to the hospital <48 hours following discharge after cardiac cath. Results: 92 patients were included in the analysis. Median age was 134 days (105-179 days) with median weight of 5.6kg (5-6.4kg). 62 patients were discharged following a 4-6 hour observation period. Of those, 3 underwent a cath intervention. Two of the 62 patients initially discharged were readmitted within 48 hours of discharge due to fever and hypoxemia. Of the remaining 30 patients, 18 stayed for 23-hour observation; 9 of those had an intervention. The other 12 patients were admitted to the hospital for >23 hours; 4 underwent intervention. There were no differences in age, weight, sex, shunt-dependence, arterial access or use of general anesthesia between those patients discharged and those admitted following cath. Patients who underwent intervention were more likely to be admitted (p<0.001), though nearly one third were discharged home without readmission. Readmission was rare (3%). No intra- or peri-procedural deaths occurred. Conclusion: Outpatient cardiac cath of infants with SVCHD can be performed with low readmission rate. Further investigation will compare cost-effectiveness of universal 23-hour overnight observation vs. outpatient discharge with potential readmission.
**#0031 PERCUTANEOUS PULMONARY VALVE IMPLANTATION WITH SAPIEN VALVES IN NATIVE AND LARGE RVOT; EARLY AND MID-TERM RESULTS**

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**Introduction:** Percutaneous pulmonary valve implantation (PPVI) has been used mainly for conduit dysfunction in right ventricular outflow tract (RVOT). Until recently, native RVOT without stenosis used to be considered a relative contraindication to transcatheter valvulation. We present early and midterm results of PPVI with Edwards–Sapien XT (ES-XT) in repaired tetralogy of Fallot (TOF) patients with native-large RVOTs.

**Method:** 53 s/p repaired TOF patients who had native RVOT with with severe/free pulmonary regurgitation, significant dilatation of the RV and without significant RVOT stenosis (peak pressure gradient between RV and main pulmonary artery (MPA) < 25 mmHg on TTE), and with a minimum RVOT / MPA diameter of ≥ 26 mm on TTE included into the study. Balloon sizing was performed with compliant (34 mm Amplatzer sizing) and semi-compliant balloons for interrogation (BI). The size of the Z-Med /BIB balloons that the Andra Stents XT would be mounted on was decided up to the indentation diameter occurred during BI; as at least 1 mm larger than the indentation diameter.

**Results:** Mean age and weight of the patients were 17 ± 7.7 (7-50) years and 49 ± 16 (22-84) kg, respectively. Before presenting pressure gradient between RV and MPA was 4.8 ± 3.4 (0-14) mmHg. Indentation diameter with BI was 26.2 ±2.7 (22-32) mm. Balloon size used for prestonenting was 28.1±2 (24-30) mm. Successful PPVI was achieved in 45 patients; 29 mm in 38 and 26 mm in seven. PPVI was performed in same session in five and 3-12 weeks after prestonenting in 40. 8 patients are waiting for valvulation after prestonenting. One patient has severe tricuspid insufficiency and underwent to surgery after valvulation. Valve function was good in all immediate after and at the last follow-up; a median of 10 months (2-25 months). RV volumes decreased and mild paravalvar leakage was observed only in five. Stent fracture has not been observed and no reintervention required yet.

**Conclusion:** PPVI with ES-XT valve, which has larger sizes as 26 and 29 mm, is feasible and safe in adolescents and adult’s patients with native RVOT without stenosis. Newer delivery systems, which is used through the smaller sheaths, gives us also an opportunity of early transcatheter valvulation in smaller patients. Prestenting for providing a secure landing zone is the most important part of the procedure. Only Andra XXL stents which has an expansion capacity up to 32 mm can be used for this purpose, currently.

**#0032 TRANSCATHETER INTERVENTIONS AFTER GLENN ANASTOMOSIS AND FONTAN OPERATION IN PATIENTS WITH UNIVENTRICULAR HEART**

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**Introduction:** Although transcatheter closure of PDA is an established standard method, most frightening complication is protrusion of the aortic disc to the DAO which may cause iatrogenic COA, especially in small children with small aorta. Ceráfex duct occluder (CDO) is a new device with similar properties with Amplatzer duct occluder (ADO). Device comes preassembled with the delivery cable by a loop connection through the holes and ready to load via the loader on the delivery cable. The loop made of surgical thread that provides the device to become flexible in 360° direction and fit to the ductal shape before releasing.

**Method:** We retrospectively evaluated 30 patients between 2007 and 2016.

**Results:** The mean age was 7.6 years (6 months-21 years) and the weight was 25.2 kg (6-54). 31 attempts were made in 30 patients. The procedures were performed after a Kawashima, Glenn and Fontan surgery in 3, 12 and 15 patients, respectively. SSD was encountered in 15 patients. Amongst these patients, closure of a Fontan fenestration was performed in 7. We occluded a decompressing vein in 5 and a pulmonary arteriovenous fistula closure in one. Closure of a residual right SVC-atrium connection was performed in one and stent implantation to reroute the hepatic blood flow to the right lung in one, after a Kawashima operation. The mean oxygen saturation of 79.3±8.1 % increased to 92.2±5.6 and the mean PA pressure increased from 11.9±2.2 mmHg (8-16) to 13.5±2.1 mmHg (10-17). LCOS and / or increased PA pressure was detected in the remaining 15. One patient was on an ECMO support. Amongst these 15 patients, an antegrade pulmonary flow was occluded using a number of devices in 7, antegrade flow was closed with the use of a covered stent, resolving an associated left PA stenosis at the same time in one. Among 4 patients suffering from branch PA stenosis, 3 received stent implantation while the remaining was treated via cutting balloon angioplasty. Two separate stents were needed to treat branch PA and extracardiac conduit stenosis in one. In the patient on ECMO support, Fontan fenestration was dilated with a balloon to ensure cardiac output at the expense of systemic desaturation. And fenestration was created in one. In patients with LCOS, the preprocedural PA pressure decreased from 20.6 mmHg (15-27) to 14.9±1.8 mmHg (11-18). There was no procedural mortality. Circulatory failure regressed in all cases except one.

**Discussion and conclusion:** To avoid reopening of the antegrade flow, surgeons should not only ligate but divide the PAs from the ventricle. In the presence of LCOS or SSD, urgent catheterization should be considered. Significant PA stenosis should be treated even if there exists no pressure gradient throughout the circulation.

**#0033 PDA CLOSURE WITH CERAFLEX OCCLUDER: IS THERE ANY ADDITIONAL BENEFIT?**

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**Introduction:** In this paper, we aimed to present transcatheter treatment of patients with a single ventricle physiology, experiencing low cardiac output (LCOS) or severe systemic desaturation (SSD) after a Glenn or a Fontan operation.

**Method:** We retrospectively evaluated 30 patients between 2007 and 2016.

**Results:** The mean age was 7.6 years (6 months-21 years) and the weight was 25.2 kg (6-54). 31 attempts were made in 30 patients. The procedures were performed after a Kawashima, Glenn and Fontan surgery in 3, 12 and 15 patients, respectively. SSD was encountered in 15 patients. Amongst these patients, closure of a Fontan fenestration was performed in 7. We occluded a decompressing vein in 5 and a pulmonary arteriovenous fistula closure in one. Closure of a residual right SVC-atrium connection was performed in one and stent implantation to reroute the hepatic blood flow to the right lung in one, after a Kawashima operation. The mean oxygen saturation of 79.3±8.1 % increased to 92.2±5.6 and the mean PA pressure increased from 11.9±2.2 mmHg (8-16) to 13.5±2.1 mmHg (10-17). LCOS and / or increased PA pressure was detected in the remaining 15. One patient was on an ECMO support. Amongst these 15 patients, an antegrade pulmonary flow was occluded using a number of devices in 7, antegrade flow was closed with the use of a covered stent, resolving an associated left PA stenosis at the same time in one. Among 4 patients suffering from branch PA stenosis, 3 received stent implantation while the remaining was treated via cutting balloon angioplasty. Two separate stents were needed to treat branch PA and extracardiac conduit stenosis in one. In the patient on ECMO support, Fontan fenestration was dilated with a balloon to ensure cardiac output at the expense of systemic desaturation. And fenestration was created in one. In patients with LCOS, the preprocedural PA pressure decreased from 20.6 mmHg (15-27) to 14.9±1.8 mmHg (11-18). There was no procedural mortality. Circulatory failure regressed in all cases except one.

**Discussion and conclusion:** To avoid reopening of the antegrade flow, surgeons should not only ligate but divide the PAs from the ventricle. In the presence of LCOS or SSD, urgent catheterization should be considered. Significant PA stenosis should be treated even if there exists no pressure gradient throughout the circulation.
Method: 21 patients underwent transcatheter closure with CDO from November 2015 to August 2016. Decision for device size selection was based on the narrowest diameter of the PDA according to the manufacturer recommendation as the aortic end of the occluder shank to be at least 1.5-2.0 mm larger than the narrowest diameter of the duct.

Results: The median age of the patients was 1.2 years (6 months to 28 years) and weight was 9.6 kg (5.4 to 82 kg). 11 patients were under one-year-old and 11 had pulmonary hypertension (mean PA pressure >25 mmHg). All patients had continuous cardiac murmur on examination and all PDAs were type A. Narrowest PDA diameter at pulmonary side was 4.1 ± 1.7 mm (2.2-8.2 mm, median 3.8 mm). Intervention was successful in all. Final angiogram showed complete closure in 17/21 of them. Echocardiography achieved complete occlusion an all on the next day. In a patient with Down syndrome PDA was closed with 4/6 mm device, and device embolized to descending aorta after persistent cough 24 hours later. Than device was snared via femoral vein approach and closed with 6/8 mm device. None of the patients showed evidence of stenosis at branch pulmonary artery and descending aorta during the follow-up.

Discussion and conclusion: Our results showed us CDO is a safe and efficacious in closure moderate to large PDAs whose duct morphology fit to the ADO I. Its uniquely designed delivery/releasing system has an advantage in view of no applying tension to the device which provides the device in stable position and not changing the device position during and immediate after the releasing. It may give us an opportunity to be sure that the device not protrudes to the aorta after releasing, especially in infants those have small descending aorta.

#0034
THE RESULTS OF EMBOLIZATION AND DISLOCATION OF THE DEVICES AFTER TRANSCATHETER CLOSURE OF ATRIAL SEPTAL DEFECT AND PATENT DUCTUS ARTERIOSUS
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Introduction: In this paper, we aimed to present the results and treatment methods of device embolization (DE) or malposition/dislocation (DD) seen after transcatheter ASD and PDA device closure. Method: Between 2004 and 2016, the patients were retrospectively analyzed regarding the DD and DE.

Results: Amongst the patients in whom a device closure of ASD was performed, DE and DD was encountered in ten and one, respectively. The median age of the patients was 19.5 (5-52) years. The septum was aneurysmatic in 2, while the rims of the defect were either deficient or thin and mobile in 5. DE occurred due to spontaneous disconnection of the devices during repeated attempts in three. The device embolized into PA in 5, into the left ventricle in 2, into the ascending aorta in one, into the right atrium in one and onto the mitral valve in one. The devices were retrieved with the use of a snare or a bioptome or a snare in 6 and the defect was closed with a larger device in 5 of these patients. 5 patients were referred for surgery. One patient was also referred for surgery due to DD at the aortic site causing significant shunt. DE and DD was encountered after PDA closure in 8. DE was observed in 5 and DD into the descending aorta (DAO) was seen in 3. The median age was 2.5 years (1 month to 8 years), and the weight was 9.5 kg (3.3-24). PDA was conical in shape in 6 and tubular in 2. The median diameter of the PDA was 8.6 mm (3.7-11.7). Except for one, pulmonary hypertension (PHT) equal to systemic pressure was present in all. Amongst 5 with DE, four devices embolized to the PA and to the DAO in one. 3 cases were referred for surgery. The devices were retrieved with the use of a snare from PA and DAO in 2. In one of these pts, PDA was occluded with a larger device. In the remaining, it could not be achieved even with a larger device. DD into the DAO led to a coarctation of the aorta in three. All of these patients had severe PHT. Two of these devices were repositioned successfully with the use of a bioptome, antegradely. In other patient, device was retrieved and PDA was closed with a muscular VSD device.

Conclusion: The risk factors for DE after ASD closure are found to be the presence of a large defect, a deficient rim with an aneurysmatic septum, mismeasurement of the defect. The most important risk factor for DE after the PDA closure are found to be the presence of severe PHT. The retrieval rate is lower after with PDA devices than ASD devices.

#0035
ECHOCARDIOGRAPHY-GUIDED TRANSCATHETER CLOSURE OF PATENT DUCTUS ARTERIOSUS IN CHILDREN: FEASIBILITY AND SAFETY OF A NEW STRATEGY
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Objective: The aim of this study was to evaluate the feasibility and safety of transcatheter closure of PDA under transesophageal echocardiographic (TEE) guidance without fluoroscopy and angiography.

Background: To avoid radiation exposure and contrast agent usage, the feasibility of transcatheter closure of atrial septal defects (ASD) without fluoroscopy has been proven. However, it is unknown that weather or not the procedure is eligible for patent ductus arteriosus (PDA) closure.

Method: From June 2014 to May 2016, a total of 102 children (38 males, 64 females), aged from 1 to 14 years (median, 2.3 years), and weighted from 6 to 46kg (median, 11.6kg), with isolated PDA (diameter of 2.5 to 7.5mm, median 3.8mm) underwent attempted transcatheter device closures. The procedures were performed under TEE guidance without fluoroscopy and angiography and the occluders were deployed by using modified delivery system via femoral venous access alone.

The patients were followed up by clinical examination, electrocardiogram, and TTE at 1, 3 and 6 months, and then yearly.

Results: PDA were successfully closed in 99 patients (97.1%). There were no acute procedural complications or severe adverse events. The procedure time ranged from 10 to 45 minutes (median, 21
minutes). Immediate complete closure of PDA was achieved in 87 patients (87.9%), and in 100% after 24 hours. The median hospital stay was 3.8 days. No intervention-related complications were detected during the period of 1-24 months (median, 12 months) follow-up.

Conclusions: Echocardiography-guided transcatheter closure of patient ductus arteriosus is a feasible and safe procedure which is eligible for alternative use in children.

Key words: Patent ductus arteriosus • Transcatheter closure • Transesophageal echocardiography • Radiation protection

#0036
RESULTS OF TRANSCATHETER PULMONARY VALVULATION IN CHILDREN < 30 KG.
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Introduction: Although widely accepted, indications for percutaneous pulmonary valve replacement (PPVr), are limited to prosthetic conduits in the right ventricle outflow tract (RVOT) in patients ≥30kg.

Aim: To evaluate the outcomes of Melody valve insertion in patients <30kg.

Methods: We analysed procedural and outcomes data from 23 patients <30 kg (9 patients <20 kg), who underwent Melody valve implantation for a native/patched RVOT(N=10), prosthetic conduit (12) and bioprosthesis (N=1).

Results: Median age and weight was 5.8 years (1.5-13) and 22kg (9-29.8). PPVr indication was regurgitation in 10, stenosis in 1 and mixed in 12. All procedures were successful. PPVr was performed through the femoral vein in 12 cases, jugular vein in 10 cases and transapical-ventricular in 1 case. Pre-stenting was performed in 95% of cases, 15/23 in the same procedure. No significant regurgitation was recorded after the procedure, and the trans-pulmonary gradient was significantly reduced. Early minor complications occurred in 2 cases (10%) The median hospital stay was 3 days (2-5). Median follow-up was 44 months (4-82). PPVr was performed through the femoral vein in 12 cases, jugular vein in 10 cases and transapical-ventricular in 1 case. Pre-stenting was performed in 95% of cases, 15/23 in the same procedure. No significant regurgitation was recorded after the procedure, and the trans-pulmonary gradient was significantly reduced. Early minor complications occurred in 2 cases (10%) The median hospital stay was 3 days (2-5). Median follow-up was 44 months (4-82). During follow-up, one patient underwent a new PPVr (valve-in-valve procedure); one patient required overdistalization of the prosthesis; and one patient developed stent fracture. Follow-up with MRI demonstrated significant improvements in right ventricular volumes and function.

Conclusion: PPVr s highly feasible in children <30 kg, in both, native RVOT and prosthetic conduits, and mid-term follow-up, demonstrates good haemodynamic results and appear promising.

#0037
STENTING OF THE NATIVE RIGHT VENTRICULAR OUTFLOW TRACT IN THE SYMPTOMATIC INFANT WITH TETRALOGY OF FALLOT (TOF)
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Objective: To assess feasibility, safety and effectiveness of right ventricular outflow tract (RVOT) stenting in symptomatic young infants with TOF.

Methods: Retrospective case note and procedure review of patients undergoing stenting of the RVOT over an 5 year period.

Results: Between 2010 and 2015, 9 patients underwent stent implantation; median age was 48 (range 1–364) days, and median weight was 3.7 (2.2–7) kg. Median procedure time was 54 (30–233) min and fluoroscopy time 20 (12.6–77) min. There was one procedural emergency surgery. Two patients required ECMO support during the procedure. The pulmonary valve was deemed unsalvageable in all patients but one (median valve diameter 5 mm (range 4–8), median Z-score -3 (range -5 to -0.5). Saturations increased from 77% (50–85%) to 95% (85–98%) (p<0.005). Two further catheter interventions were undertaken (balloon in 1, stent in 1). 7 patients underwent delayed surgery (complete repair in all of them) at a median of 140 (111–200) days post-stenting. No surgery was compromised by the presence of stents in the RVOT. There were no perioperative deaths. Median Nakata index increased from 80mm2/m² (50–200) to 126 mm2/m2 (100–251)(p <0.05) before surgical repair.

Conclusions: In the symptomatic young infant with TOF, stenting of the RVOT provides a safe and effective management strategy, improving arterial oxygen saturation and encouraging pulmonary artery growth.

#0038
PERSISTENT 5TH AORTIC ARCH IN A PATIENT WITH 22Q11.21 DELETION
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Background: Double-barreled aorta or persistence of 5th aortic arch is an extremely rare cardiovascular anomaly and only few cases of persistence of 5th arch with Pulmonary atresia, ventricular septal defect, and aorto-pulmonary collaterals (PA, VSD, MAPCA) have been reported. We present a case of PA, VSD, MAPCA with persistence of the 5th aortic arch in a patient with a 22q11.21 mutation. This double barrel aorta is characterized by presence of a second systemic to systemic channel connecting the ascending and descending components of the aorta.

Case Report: A full term female newborn failed pulse oximetry screening at 36 hours of life. First echocardiogram showed PA, VSD, MAPCA and discontinuous pulmonary arteries. Prostaglandin drip was started and patient transferred to our institution. Repeat echo followed by CT angiography confirmed anatomy and showed left sided arch with a persistent 5th arch.

In our patient a second vessel originated from the undersurface of the proximal transverse aortic arch (embryologic 4th aortic arch) just distal to the origin of the right common carotid artery and connected to the distal transverse aortic arch just distal to the origin of the left subclavian artery. The normal (4th) arch is small compared to the 5th arch and sequentially supplies the left common carotid, left vertebral, and left subclavian arteries before connecting to the 5th arch. The left sided ductus originated distally from this connection from the
**Background:** This study aimed to evaluate whether an ultrasound-guided access technique can improve upon the landmark-guided technique in achieving quicker vascular access in the femoral artery and vein during pediatric cardiac catheterization.

**Methods:** A random sample of patients greater than 60 days of age (Group 1) and under 60 days of age (Group 2) were chosen during a time period (before 2013) when access was obtained using only landmark-guided techniques and during a time period (after 2013) when ultrasound-guidance was used. Data collected included age (years), height (cm), weight (kg), body surface area (m²), access site/s, sheath size, case times, time of first sheath entry, time of final sheath entry, and total time to obtain access (minutes).

**Results:** Data was collected on a total of 101 patients for Group 1 and 50 patients for Group 2 in each time period. Patients were matched in regards to height, weight, and body surface area between the two time periods.

Additionally, the reimbursement benefit over a 6 month period in 2014 was $30,000.

**Conclusion:** Our findings show that the use of an ultrasound-guided technique to access the femoral artery and vein significantly decreases the time to vascular access and has beneficial financial implications.

<table>
<thead>
<tr>
<th></th>
<th>Time to 1st Sheath Entry P</th>
<th>Time to Final Sheath Entry P</th>
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<tbody>
<tr>
<td>Group 1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pre U/S (n=101)</td>
<td>6.6 (±7.1)</td>
<td>15.0 (±13.0)</td>
</tr>
<tr>
<td>Post U/S (n=101)</td>
<td>5.9 (±5.8)</td>
<td>11.9 (±10.5)</td>
</tr>
<tr>
<td>Group 2</td>
<td></td>
<td></td>
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<tr>
<td>Pre U/S (n=50)</td>
<td>12.1 (±13.5)</td>
<td>25.9 (±20.3)</td>
</tr>
<tr>
<td>Post U/S (n=50)</td>
<td>8.7 (±8.1)</td>
<td>17.2 (±13.4)</td>
</tr>
</tbody>
</table>

Data is presented as mean (±standard deviation).

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**#0038**

**EVALUATION OF RADIATION EXPOSURE AND WEIGHT VARIABILITY DURING CARDIAC CATHETERIZATION FOR CONGENITAL HEART DISEASE: DATA FROM A MULTICENTER BRAZILIAN REGISTRY**

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**Introduction:** Cardiac catheterization continues to be of great value in the field of congenital heart disease. There is an increasing concern of radiation dosage in the last years because of its known deleterious

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**#0039**

**BENEFICIAL EFFECT OF ULTRASOUND-GUIDED VASCULAR ACCESS IN THE PEDIATRIC CARDIAC CATHETERIZATION LAB**

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Figure 1 (#0038). Still frame of descending aorta angiogram depicting the double barrel aorta.

...undersurface of the distal aortic arch and supplied a tortuous Aorto-pulmonary collateral (APC) to the right lung. The left lung was supplied by one large APC. (Fig 1)

Patient underwent stenting of the ductus and came off prostaglandin and was discharged home. Hypocalcemia was noticed during intensive care stay. Chromosomal microarray revealed a deletion of 2.54 MB on chromosome 22q11.21 involving several genes including the TBX1 gene.

**Discussion:** As per our knowledge this is the first case report which shows the association of persistence of 5th aortic arch with PA, VSD, MAPCA and TBX1 gene deletion. TBX1 gene deletion has been shown to cause dysfunction of the neural crest cell and anterior heart field leading to the various cardiovascular anomalies in 22q11 del syndromes.

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**#0040**

**EVALUATION OF RADIATION EXPOSURE AND WEIGHT VARIABILITY DURING CARDIAC CATHETERIZATION FOR CONGENITAL HEART DISEASE: DATA FROM A MULTICENTER BRAZILIAN REGISTRY**

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²Instituto Dante Pazzanese, São Paulo, Brazil

**Introduction:** Cardiac catheterization continues to be of great value in the field of congenital heart disease. There is an increasing concern of radiation dosage in the last years because of its known deleterious
effect. Children have more radio sensitivity but the evaluation of this exposition is more difficult mainly due to the weight variability. Finally, radiation dosage is not yet well standardized in this population.

Methods: This was a multicenter observational study. Data was collected from a Brazilian Registry of cardiac catheterization on congenital heart disease from March 2013 to June 2014. Patients aged more than 18 years old were excluded from this study. Four hundred and ninety five patients had dose-area product (DAP) measured and were included in this study. Median and interquartile intervals (25th and 75th) of age and weight were respectively 50 months (10 ; 103) and 15 Kg (7 ; 28).

Results: The median and interquartile intervals (25th and 75th) of DAP was 742.2 (288.8 ; 1791.5) uGy.m2. Interventional procedures had higher DAP than diagnostic procedures (25th, 50th, 75th percentiles: 230, 715, 1534 versus 315, 751, 2095 uGy.m2). DAP correlated with body weight \( (r = 0.56) \) but best correlated with weight-fluoroscopic time product \( (R = 0.75) \). DAP/weight showed no difference between diagnostic and interventional procedures \( (25th, 50th, 75th \text{ percentiles: 23, 57, 110 versus 30, 57, 139}) \).

Conclusions: DAP/weight has been considered in the last few years as the most reliable unit to evaluate radiation exposure in pediatric population. However, to date, there are only two reports in the literature. This study reports similar doses when comparing to the benchmark doses previously published by the CCISC. It is very useful to report these data in order to be continuously evaluating the impact of strategies to lower radiation exposure in this vulnerable population. Differences between institutions can alert for increased radiation exposure in patients and staff and support a program of quality improvements for radiation safety.

#0041

HYBRID BALLOON VALVULOPLASTY FOR PULMONARY ATRESIA AND INTACT VENTRICULAR SEPTUM: 10 YEAR EXPERIENCES

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Objective: Management of pulmonary atresia with intact ventricular septum (PA/IVS) remains challenging for the unfavourable outcomes after surgical or percutaneous interventional approaches. This study aimed to report outcomes of our hybrid balloon valvuloplasty via sternotomy for PA/IVS.

Methods: From 2005 to 2015, consecutive patients with PA/IVS who underwent hybrid balloon valvuloplasty in our institution were included into the current study. Ductal ligation, bidirectional Glenn shunt (BDG) and modified B-T shunt (mBT) were applied according to the post-valvuloplasty hemodynamics. Exclusion criteria included: right ventricular (RV) dependent coronary circulation and muscular atresia.

Results: A total of 77 consecutive patients were recruited. The median age at balloon valvuloplasty was 3 months (1 day to 24 months). Fifteen (19.4%) patients were neonate, 19 (24.7%) patients had mild or moderate RV hypoplasia, and more than moderate tricuspid regurgitation presented in 65 (84.1%) patients. Procedures included 25 valvuloplasty alone, 27 valvuloplasty + ductal ligation, 20 valvuloplasty + ductal ligation + mBT, and 5 valvuloplasty + ductal ligation + BDG. RV outflow tract continuity were established in all patients without early mortality and early reoperation. After valvuloplasty, SO2 increased from 73.5 ± 25.7% to 93.0 ± 3.4%. Four (5.2%) patients had a residual pressure gradient more than 30 mmHg. During 6.5 years (4 months-11 years) follow-up, there were 7 late deaths, 2 re-balloon dilatations, 2 ductal occlusions and 2 mBT occlusions. According to the hemodynamics, 2 patients underwent subsequent BDG and 2 patients underwent subsequent total cavopulmonary connection. At the latest follow-up, 59 patients had biventricular circulation, 5 patients had one and a half ventricular circulation and 2 patients had univentricular palliation. In multi-variate analysis, tricuspid z score was identified as independent protective factor for post-operative death \( (p=0.02, OR=0.21) \).

Conclusion: Outcomes of hybrid balloon valvuloplasty for PA/IVS is favorable with satisfactory potential to biventricular circulation.

#0042

NOVEL TRANSCATHETER INTERVENTION IN COR TRIATRIATUM DEXTER

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Cor triatriatum Dexter is an extremely rare congenital anomaly (0.025%), in which the right atrium is divided into two chambers by a septum, diagnosed on autopsy in the past. We describe ante-mortem diagnosis, by 2D transthoracic echocardiography with agitated saline contrast echocardiography and inferior venacava or superior venacaval venography. To the best of our knowledge, for the first time in world, we report a novel method of transcatheter balloon disruption of membrane in five cases, along with the balloon dilatation of rheumatic mitral stenosis in two cases and device closure of atrial septal defect in one case, to prevent morbidity and mortality.

Objective: To describe the importance of transthoracic echocardiography with agitated saline contrast echocardiography and assess the feasibility and efficacy of transcatheter intervention in Cor triatriatum Dexter.

Material and Results: Out of fifteen consecutive patients of Cor triatriatum Dexter diagnosed with transthoracic echocardiography with agitated saline contrast echocardiography, five patients who underwent transcatheter balloon disruption of membrane and other interventions formed the material for study. Three patients were boys and two were girls, age ranged between 3 to 17 years, median age was 10 years. Three patients presented with exertional dyspnea and two were asymptomatic. Two who had rheumatic heart disease with mitral stenosis underwent balloon mitral valvuloplasty and one case with atrial septal defect without pulmonary artery hypertension underwent device closure.

Conclusion: The Cor triatriatum Dexter is not benign as mortality occurs due to pulmonary embolism. Timely diagnosis with transthoracic echocardiography with simple agitated saline contrast echo followed by balloon disruption can prevent cyanosis, pulmonary artery hypertension, morbidity and mortality.
#0043
CHALLENGES OF TRANSCATHETER INTERVENTIONS FOR CONGENITAL HEART DISEASES IN DEXTROCARDIA
I.B Vijayalakshmi
BMC&RI Super Specialty Hospital (PMSSY), Bengaluru, India

Background: Several challenges are faced by interventionists while performing various percutaneous interventions for congenital heart disease (CHD) in patients with dextrocardia. The anatomical alterations in dextrocardia especially the lie of the interventricular septum (IVS) can cause impediment for device closure of ventricular septal defect (VSD) and lie of interatrial septum (IAS) for puncture.

Aim: The aim of our study is to evaluate the challenges, feasibility and efficacy of transcatheter interventions in children with CHD in dextrocardia.

Materials and Results: Out of 60 patients of CHD with dextrocardia catheterized, only 9 patients (15%) underwent transcatheter interventions. The age was 4 months to 16 years (mean 5.4 years), weight - 4.1 to 40 kgs (mean 8.4 kgs). 3 cases underwent successful device closure for patent ductus arteriosus (PDA), 2 cases of midmuscular VSD (MVSD) were closed. One had to be closed with Amplatzer ventricular septal occlude through the jugular approach and the other with Amplatzer duct occluder II (ADO II). There was a difficulty in puncturing IAS during balloon valvuoplasty for mitral stenosis in a case of right-sided May Thurner Syndrome (MTS). Balloon valvuoplasty was done in one infant with severe pulmonary stenosis by flipping the cine image. One very sick patient with inferior vena cava web died after cavoplastry and stenting.

Conclusion: The catheter interventions in CHD with dextrocardia though difficult is feasible. The device closure of PDA and MVSD is not difficult especially with ADO II. The balloon mitral and aortic valvuloplasty in the complex cardiac anatomy of situs inversus totalis is feasible and safe. Rarely right-sided MTS may cause problem for right femoral access during transcatheter procedure.

#0044
CHALLENGES OF INTERVENTIONS FOR ASSOCIATED LESIONS IN CASES OF APICAL NON-COMPACtion
I.B Vijayalakshmi
BMC&RI Super Specialty Hospital (PMSSY), Bengaluru, India

Background: Isolated left ventricular non-compaction is reported extensively. But apical non-compaction (ANC) of both ventricles and septum is not reported much in literature. For the first time in the world, we are reporting the challenges of various interventions for different associated lesions in ANC.

Aim: To know the challenges and feasibility of transcatheter interventions for the associated lesions in cases of ANC to reduce the pump failure.

Material and Results: Out of 100 consecutive patients diagnosed as ANC by transthoracic echocardiography (TTE), 30 cases underwent various transcatheter interventions, formed the material for this study. Age ranged 3 days to 17 years (mean 8 years). The device closure was done for PDA in 2, VSD in 15, ASD -1, ARVT in 1, ARAT in 1, ABV in 4, PBV in 2, aortoplasty in 1, PTMC in 2, pericardiocentesis in 3. 5 patients underwent two procedures in the same sitting. They were ABV and PBV, ABV and PTMC, ABV and PDA device closure, ASD and VSD device closure and PDA and VSD device closure. 3 cases of VSD were post-operative residual and one was closed with multiple devices. One 8 months infant had apical VSD closed with ADO II. Another 2 year old underwent hybrid surgery for closure of VSD with 14 mm device. One child with mirror image dextrocardia and midmuscular VSD was closed with device. In one case procedure was abandoned as 18 mm VSD device slipped.

Discussion: Procedures in ANC is risky in presence LV/or RV dysfunction with or without thrombosis. Positioning the device in apical VSD in ANC cases is very challenging as the device gets caught in trabeculae in RV and if more tug is given the device slips through spongy myocardium. The results of interventions are very gratifying as the superadded pump failure due to pressure or volume overload caused by associated lesions improves significantly. One patient with severe AS and mitral stenosis had reverse May Thurner syndrome (obstruction of right common iliac vein by right common iliac artery), hence procedure was done through left femoral puncture.

Conclusion: Associated lesions in ANC worsen the pump failure. Transcatheter interventions though challenging are feasible safe effective and are lifesaving. Transcatheter interventions certainly reduce the morbidity and mortality in ANC patients who are at high risk for surgery or redo surgery.

#0045
RETROGRADE TRANSCATHETER CLOSURE OF VENTRICULAR SEPTAL DEFECTS
I.B Vijayalakshmi
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Background: Amplatzer Duct Occluder II (ADO II) is designed for closing long ducts in infants. There are few reports of “off-label” use of ADO II in non-ductal positions.

Aim: To evaluate the advantages and disadvantages of retrograde transcatheter closure of ventricular septal defects (VSD) with ADO II.

Material and Results: 102 cases of VSDS closed by retrograde transcatheter method with ADO II, formed the material for the prospective study. Age: 8 months to 23 years (mean 9.1 years). 74 perimembranous VSDs, 14 muscular VSDs, 13 Gerbode defects, one midmuscular VSD with dextrocardia, were closed. The shortest fluoroscopic time was 4.2 min, mean was 8.4± 4.1min. In six cases there was initially a small residual shunt which had closed on three months follow up. Only in one case the device embozized to left pulmonary artery and it was retrieved . Eleven cases developed transient complete heart block which resolved and only one of them needed temporary pacing.

Discussion: ADO II has a very low profile and can be easily delivered through a 5F guiding catheter needs very short fluoroscopic time as artero-venous (AV) loop is not needed. The cost is 1/3 the cost of
regular ventricular septal occluder. However, it is not useful in VSDs measuring more than 6 mm and in those with insufficient aortic rim.

Conclusion: ADO II is an excellent device in ventricular septal defects. The procedure time and the cost are significantly lesser than regular devices. The success rate is very high and complication rate is very low.

**#0046**
TRANSCATHETER DUCTUS ARTERIOSUS STENTING WITH EXPECTED REDILATION AS A SURGICAL ALTERNATIVE
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**Background:** Transcatheter ductus arteriosus stenting (DAS) has been used to provide a reliable temporary source for pulmonary blood flow in cyanotic congenital heart patients. However, even in the context of surgical mortality rates approaching 10%, most modern congenital cardiovascular centers continue to rely almost exclusively on surgical shunts. Data on DAS from centers with sophisticated surgical services is limited. We report DAS experience from a modern comprehensive congenital heart center in North America.

**Methods:** This is a single-center retrospective review of all patients who were taken to the cardiac catheterization lab with the intent of DAS from January 2006 to August 2016.

**Results:** Successful DAS was performed in 30/32 patients (97%), median age 7 days (range 1-176), median weight 3.43kg (1.5-6.8). Diagnoses included 14 PA/IVS, 5 critical PS, 4 TOF, 2 PA/VSD, 3 TGA, 1 DORV/TGA, 1 tricuspid atresia, 1 Ebstein’s & 1 Heterotaxy/AVC/PS. 2/32 required urgent surgical shunt placement, due to ductal spasm prior to stenting in one and stent thrombosis in the other. Bare metal coronary artery stents were used, median stent diameter 3.5 mm (2.5-4). Median saturation at discharge was 91% (80-99). At a median follow up of 3.9yrs (2 mos-9.4yrs), 17 (55%) had reinterventions for low saturations (balloon dilation alone n=5 and restent n=12). Median time between implant and re-intervention was 74 days (33-186). Stent thrombosis at the time of re-intervention occurred in 1/17; patient was referred for an early Glenn. Complications involving femoral vessel compromise occurred in 7 (22%), 3 resolved with anticoagulation, 4 formed sufficient collaterals. 16/30 progressed to surgical repair or next stage palliation; 14 did not require further surgical intervention with adequate prograde pulmonary blood flow (balloon pulmonary valvuloplasty n=13), bilateral DA stenting as destination palliation (n=1). There were no deaths associated with the DAS. One patient died after a Glenn procedure while awaiting heart transplant 3 months post DAS.

**Conclusions:** DAS appears to be a safe and effective procedure for establishing a reliable temporary source of pulmonary blood flow in selected cyanotic congenital heart patients. Reintervention for stent redilation is common and should be a planned component of this management strategy. It is likely that more experience from centers with sophisticated surgical services will be necessary before community practice evolves toward greater reliance upon DAS.

**#0047**
RE-STENTING IMPROVES VESSEL PATENCY, LUMEN AREA AND ENDOTHELIALIZATION: A FEASIBILITY STUDY IN PIGLETS DEMONSTRATING RE-STENTING WITH SIMULTANEOUS INTENTIONAL FRACTURE OF PREVIOUSLY PLACED STENTS
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**Background:** Intentional stent fracture in-vivo has shown medial dissection/vessel injury. Spontaneous stent fracture in humans has been reported to cause stent collapse, hemodynamic compromise and embolization of stent fragments, many of which were treated with re-stenting.

**Objective:** To demonstrate feasibility and propose re-stenting prior to intentional stent fracture to avoid such complications. We sought to compare stent fracture with and without a stabilizing additional stent and study the effects on the tissue.

**Methods:** Five months after fourteen low profile stents where implanted in the aorta of four piglets, they were intentionally fractured using high-pressure balloons, one group with another stent placed inside (re-stent group) and one without (single stent group). Vessel/stent were studied at both short-term and long term intervals. Results: The re-stent group demonstrated significantly larger vessel segment (113±28mm² versus 59±17mm², p=0.0476), less long term luminal diameter loss (45.5% versus 73.7%, p=0.0065), lack of strut protrusion and higher degree of endothelization (95.7±2.1% versus 74.1±7%, p=0.0145). There was similar degree of vessel wall injury at the time of the fracture, however at long term evaluation the injury score in the re-stent group demonstrated statistically significant improvement (p=0.0426). No damage to the external part of the blood vessels or the surrounding soft tissue was noted in either group.

**Conclusion:** Re-stenting at the time of intentional stent fracture provides numerous advantages including larger vessel diameter, maintained vessel patency, more complete endothelization, lack of strut protrusion with decreased chance of injury and future complications. Human studies are needed to confirm our observations.

**#0048**
VSD CLOSURE USING Cera® DEVICES
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**Objective:** The aim of this study is to describe the immediate to long follow up in 15 patients (p) with perimembranous (PMVSD)
Background: Cera VSD Occluder is formed of a nitinol wire mesh, covered with a ceramic coating which reportedly reduces the risk of thrombus formation, encourages endothelialization, and minimizes systemic nickel ion release.

Material and Method: from October 2013 to September 2016, 14 p were treated with CERAO devices. This is a prospective analysis. Median age: 11 y/o (2.6 to 25 y/o). Median weight: 40 kg (14 to 70 kg). Gender: 9/14p M 6/14p F. All the p had femoral artery and venous approach. All the p with general anesthesia: X: QP/QS: >1.3:1. All the procedures were carried out with arterio-venous loop. LV angiograms: a) perimembranous VSD: 60° cranial-20° LAO, b) muscular VSD: 40° cranial-20° LAO. All the procedures were guided with TEE and TTE. Measurements by echo and angiography: right and left diameter and length of the defect.

Results: 7/14 p: perimembranous VSD, 3/14 p post surgical (1/14 p double committed VSD, 1p PM VSD and 1/14 p Pulmonary atresia with VSD), 4/14 p: muscular VSD (1/14 mid ventricular VSD, 1/14 multiple VSD, 2/14 muscular-membranous VSD). Total procedures: 17 (1 p suffered an AV block during the catheterization of the VSD so that the procedure was cancel and in a second session the VSD was totally occluded). 1p the device embolized. In a second procedure, the device was retrieved and a bigger device was successfully positioned. Devices used: 1 type I, 8 type II, 2 type III, 3 muscular VSD. Complications: 1 embolization, the device was snared and the VSD was treated with a bigger device. This embolization was in the third p. 3 AV blocks in the first 24 hs, treated with steroids. The p were discharged with sinus rithm. One of them had sick sinus syndrome.

Follow up: 3p with residual shunt (2 mild and 1 moderate who is waiting for a second procedure). There weren’t av-block. As the initial experience was good, we decided to close VSDs in p under 10 kg. 1p with 8 grs had a perimemb VSD. It was closed with 7 type III perimemb device with correct result.

Conclusions: 1) Lifetech devices are a good choice to close VSDs. 2) The AV-blocks after procedures were considered as inflammatory process for the maneuvers. The steroids diminished the inflammation. 3) The embolization described was due to the little experience. 4) There weren’t AV-blocks during follow up.

Methods and Results: 31 catheterizations performed on 25 patients (18 males) weighing <10kg from Jan 2008 to June 2016 with the intention to relieve RVOTO were retrospectively reviewed. 18 procedures were first palliations, 2 with initial surgical palliation, 6 being restenting procedures and 5 were on post-operative RVOTs. The primary diagnosis was Tetralogy of Fallot including double outlet ventricle-arterial connection in 10.9 p patients (36%) had associated cardiac lesions, with atroventricular septal defect being the commonest (n=4, 16%). Prematurity and genetic disorder accounted for 32% and 24% respectively. The indication for RVOT stenting were low body weight precluding surgery (n=16, median 2.59kg, IQR 1.9-3.42kg), high risk associated cardiac lesions (n=5), high risk comorbidities (n=3), severe cyanotic spells (n=2) and post-operative residual RVOTO (n=5). 9 (36%) patients were on prostaglandin infusion, 9 (36%) ventilated and 3 (12%) were on propranolol. Eight procedures (26%) were performed as emergent or salvage therapy. Overall procedural success rate was 93.5% (29/31 procedures) with 32 stents implanted and 3 procedures required 2 stents. Significant improvement of SaO2 was observed (71±14.6% to 88.7±7.9%; p<0.001). Six complications were recorded (19%) including 1 cardiac perforation that required no treatment, and 1 sudden cardiac arrest during diagnostic catheterization, the only in-lab mortality; others were minor complications. Catheter reinterventions were necessary for 11 patients, 2 patients required 2 reinterventions. Median time to reintervention was 8.4 months (IQR=0.8-14.5). A lower main pulmonary artery z score (-4.3±1.7 vs -2.4±1.4, p=0.02) and pulmonary annular z score (-5.2±1.5 vs -3.4±2.2, p=0.04) were associated with reintervention. Twenty patients survived definitive surgery, with a mean RVOT velocity of 2.6±0.7m/s from their latest echocardiogram; while 2 patients were awaiting definitive surgery.

Conclusion: RVOT stenting achieved good palliation with high success rate in this group of high-risk symptomatic TOF patients.

#0050
SUCCESSFUL TRANSFEMORAL TREATMENT OF AN INNOMINATE ARTERY PSEUDOANEURYSM COMPLICATION FROM EXTRACORPOREAL MEMBRANE OXYGENATION IN A NEONATE WITH REVIEW OF THE LITERATURE

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Background: We report successful repair of a large pseudo-aneurysm in a 7 week neonate which developed off of the innominate artery as a complication of ECMO cannulation. From review of the literature, this is the first reported case to be successfully treated via transfemoral approach using covered stents.

Methods: A full term infant with history of biventricular hypertrophy and pulmonary hypertension was placed on VA ECMO soon after birth. At 7 weeks of life, post decannulation, a large pseudoaneurysm was found off of the innominate artery by echocardiogram. A CT with contrast confirmed the diagnosis. During cardiac catheterization, angiography revealed that the pseudoaneurysm measured 24 mm x 25 mm. The diameter of the subclavian artery was 4 mm and the origin of the innominate artery measured approximately 5mm. The length that needed to be covered was approximately 25 mm. A wire was placed in the distal subclavian artery and exchanged of a 6F x 55 cm Flexor...
Tuohy-Borst delivery sheath. We deployed two 6 mm x 22 mm Atrium Cast premounted covered stents in a tandem fashion to effectively isolate the aneurysm and maintain flow to the subclavian artery. Result: Follow up angiography demonstrated excellent result with nearly complete occlusion of the pseudoaneurysm and normal opacification of the right subclavian artery through the stent. There were no procedural or post procedural complications. Patient was electively started on heparin after sheath removal for 6 hour. The femoral artery pulses remained normal. Follow up echocardiography revealed trace shunt from the where the stents were overlapped. The patient was transferred to the NICU and discharged home when his condition stabilized.

Conclusion: We reviewed the literature and found one case report where pseudoaneurysm had developed after ECMO, but no account was made as to the management of the pseudoaneurysm. Hence, to our knowledge, this is the first successfully closed pseudoaneurysm case in a neonate. Management options in this patient included transcatheter management or proximal ligation of the innominate artery to prevent further dilation and rupture. Since the patient had the right carotid artery ligated after the ECMO cannula was removed, we wanted to maintain adequate blood flow to the right upper extremity and avoid steal phenomenon form the vertebral artery.

#0051 PERCUTANEOUS STABILIZATION OF DUCTAL-DEPENDENT PULMONARY BLOOD FLOW USING PDA STENT IS ASSOCIATED WITH DECREASED MORBIDITY COMPARED TO SURGICAL SHUNT PLACEMENT

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Background: Transcatheter stenting of the patent ductus arteriosus (PDA) is becoming an accepted alternative to surgical aortopulmonary shunt for stabilization of ductal-dependent pulmonary blood flow. However, there is limited data comparing procedural and intermediate-term morbidity between these two palliative strategies. Methods: This is a retrospective single-center review of patients who underwent stenting of the PDA or surgical aortopulmonary shunt from 2005 to 2015. Primary outcomes were survival, length of stay and freedom from reintervention. Patient charts and the Society for Thoracic Surgeons database were queried for procedure-related adverse events. Results: 55 patients were referred for PDA stenting and 55 patients were referred for surgical intervention with a Blalock-Taussig or central shunt. PDA stent placement was successful in 44 (80%) patients. Baseline characteristics were similar between the 2 groups. While there was no significant difference in 30-day survival (95.4% vs. 97.9%, p=0.24), there was a trend towards improved overall survival in the PDA stent group (93% vs. 81.1%, p=0.09, median follow-up 941 vs. 1697 days, respectively). The proportion of patients who underwent biventricular repair was similar between study groups. Patients with PDA stents had shorter median ICU and hospital length of stays (2 vs. 7 days, p=0.04; 6 vs. 17 days, p=0.01, respectively), were less likely to require nasogastric tube feeds at discharge (p<0.01) and were less likely to require invasive intervention in the first 30 days after initial palliation (p=0.04). There were more adverse events in the surgical shunt cohort (119 vs. 36, p<0.01), although the proportion of patients who experienced ≥1 adverse outcome was not different (67.3% vs 50%, p=0.08). The surgical cohort had a greater incidence of mechanical ventilation requirement ≥7 days (p=0.02), wound infection (p=0.01), unplanned cardiac catheterization (p=0.05), surgical exploration (p=0.04), and low cardiac output syndrome (p=0.02). The incidence of venous or arterial thrombosis in the PDA stent cohort was relatively low (6.8% & 11.4 %, respectively). CONCLUSION: Percutaneous stenting of the PDA is an effective strategy for stabilization of ductal dependent pulmonary blood flow. While there were no significant differences in early and overall mortality between cohorts, PDA stenting was associated with significantly less morbidity and need for re-intervention.

#0052 INFERIOR VENA CAVA COMPLETE OCCLUSION AFTER LIVER TRANSPLANT TREATED WITH RADIOFREQUENCY PERFORATION AND STENTING IN A PEDIATRIC PATIENT.

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We present a 15 month old boy with biliary atresia, who underwent liver transplantation. Surgery was complicated by a bowel perforation and thrombosis of the hepatic artery. Four weeks after the incident he was retransplanted. A thrombosis at the inferior vena cava anastomosis was diagnosed by ultrasound in the immediate post-surgical period. Five months after surgery an increase in serum aminotransferases (8 x normal values) and gamma-glutamyl trasferases (2 x normal values) was observed. A liver biopsy showed signs of hepatic vein outflow obstruction. A cavography showed the complete occlusion of the inferior vena cava with partial decompression towards the ayzygos vein system. A 16.9 mm distance between superior and inferior stumps of the IVC was measured. After multidisciplinary discussion, the patient was brought to the catheterisation laboratory, where a radiofrequency needle (Baylis medical, Toronto, Canada) was advanced via the inferior stump, reaching the right atrial stump of IVC. The guidewire was snared and exteriorized by the right jugular vein. On that guidewire rail, a 28 mm covered CP stent (NuMed, Hopkinton, USA) was deployed using a 10 mm x 3 cm BIB balloon (NuMed). The initial clinical response was very encouraging, with ascites significantly reduced and normalisation of laboratory tests. The patient needed balloon redilations, due to caudal stenosis of the stent, at two and four months post-procedure. One year after the procedure the stent is still permeable and without significant gradient. The patient is still under enoxaparin and aspirin. Liver function is normal.

Conclusion: Percutaneous approach of this rare complication was the alternative to avoid a third liver transplant in our patient. The good results and the feasibility of the technique make it a very encouraging option in this group of patients.

#0053 THE EFFECT ON SOMATIC GROWTH OF SURGICAL AND TRANSCATHETER TREATMENT OF SECUNDUM ATRIAL SEPTAL DEFECTS

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EXCIMER LASER THERAPY FOR NEONATAL CONGENITAL HEART DEFECTS WITH CRITICAL OBSTRUCTION: A SINGLE INSTITUTIONAL EXPERIENCE.
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Interventional procedures for neonatal congenital heart defects with critical obstruction have evolved over the past 25 years with variable results. The excimer laser catheter (Spectranetics, Corporation) is an over the wire system that is available in 4 diameters (0.9mm; 1.2mm; 1.5mm and 2.0mm) and creates a distinct circumferential orifice equivalent to the catheter diameter at the site of intervention.

Methods: Since 2001, we have utilized the Excimer laser system, with or without a guide catheter, for the treatment of neonatal congenital heart defects with critical obstruction vs other various established methods including surgical intervention.

Results: Ten neonates with pulmonary valve atresia/critical stenosis and intact ventricular septum, 4 neonates with hypoplastic left heart syndrome and intact atrial septum, 1 premature neonate with an occluded right pulmonary artery secondary to thrombus and 1 Fontan baffle with failing physiology underwent successful perforation and additional intervention using the excimer laser system (videos of the technique would be provided for oral presentation). All laser perforations were successfully performed utilizing fluoroscopy with or without transesophageal echocardiographic imaging. In the case of pulmonary valve atresia and HLHS, only the 0.9mm diameter laser was utilized which facilitated without difficulty, the ability to dilate the valve or atrial septum with a single large diameter Mini-Tyshak balloon thereby avoiding the need for a gradational approach. The single neonate with an acutely occluded right pulmonary artery underwent serial laser dilation with all 4 diameter laser catheters prior to conventional balloon angioplasty. There was a single inadvertent perforation of the RV outflow tract which sealed spontaneously and no other complications were reported secondary to the laser procedure.

Conclusion: Excimer laser therapy is an excellent alternative therapy for the treatment of neonatal congenital heart disease with critical obstruction. The creation of a distinct 1mm or greater orifice at the site of intervention, facilitates the utilization of a single diameter balloon dilation catheter, thereby avoiding a gradational approach and reducing fluoroscopy exposure.

EXPERIENCE.
Methods: Retrospective chart review of patients post-SPVR or TPVR at Cohen Children’s Medical center of New York from 2013-2015. Volumetric data obtained from cardiac magnetic resonance (CMR), one year prior and about one year after PVR. PS was defined as peak-peak gradient ≥35mmHg by catheterization or peak gradient ≥50mmHg by echocardiography. Student’s t-test and Mann-Whitney test for group comparisons and chi-square and Fisher’s exact test for associations.

Results: 30 patients post-PVR: 15 surgical, 15 transcatheter (1 hybrid). At referral, the SPVR group had only PR; indications for SPVR were: 2+ CMR parameters in 12 patients; 3 had decreased left ventricular ejection fraction (LVEF) and hemodynamic findings and/or abnormal exercise stress test. In the TPVR group, 8 patients had PR, 1 had PS and 6 with PR/PS; indications for TPVR were: 2 CMR parameters in 5 patients, 10 had symptoms and/or hemodynamic findings. At baseline, SPVR group had significantly larger right ventricular (RV) volumes than TPVR group. Biventricular function was not significantly different. Post-PVR, both groups had significantly decreased RV volumes and increased LV diastolic volumes. The SPVR group improved LV cardiac output (CO) (4.9±1.1 to 5.9±0.8 L/min, p=0.03) and biventricular function (RVEF 44.7±5.4% to 53.4±8.8%, p=0.02, LVEF 54.2±5.4% to 60.2±3.8%, p=0.04) whereas the TPVR group had no significant changes.

Conclusions: Most patients were referred for SPVR due to CMR volumetric criteria, whereas for TPVR due to exercise intolerance with mildly abnormal CMR criteria. One year after PVR, both groups had near-normal biventricular volumes and function irrespective of baseline characteristics at referral.

#0056
COMPARISON OF PATIENTS UNDERGOING SURGICAL VERSUS TRANSCATHETER PULMONARY VALVE REPLACEMENT: CRITERIA FOR REFERRAL AND MID-TERM OUTCOME
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Background: Pulmonary regurgitation (PR) and/or stenosis (PS) is challenging in patients with congenital heart defects. We compared baseline characteristics of patients undergoing surgical (SPVR) versus transcatheter (TPVR) pulmonary valve replacement and identified if criteria for referral differ.

Cases: A 23 year old with bicuspid aortic valve and supra valvar aortic narrowing received a Palmaz-XL 3110 stent placed at 22mm. A 35 year old with Tetralogy of Fallot with supra valvar narrowing due to bypass cannulation received a 26mm EV3 placed at 25mm. An 11 year old post-transplant (cardiomyopathy) received a 36mm EV3 at 24mm. There was resolution of gradient, angiographic improvement, and no complications in all cases.

Conclusion: Our experience suggests that transcatheter ascending aorta stenting is a safe and feasible alternative to surgery in selected cases. 3D Rotational Angiography provides an optimal roadmap for successful implantation.

#0055
COMPARISON OF PATIENTS UNDERGOING SURGICAL VERSUS TRANSCATHETER PULMONARY VALVE REPLACEMENT: CRITERIA FOR REFERRAL AND MID-TERM OUTCOME
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2Children’s Hospital of Colorado, Denver, Colorado, USA

Background: Pulmonary regurgitation (PR) and/or stenosis (PS) is challenging in patients with congenital heart defects. We compared baseline characteristics of patients undergoing surgical (SPVR) versus transcatheter (TPVR) pulmonary valve replacement and identified if criteria for referral differ.

Methods: Retrospective chart review of patients post-SPVR or TPVR at Cohen Children’s Medical center of New York from 2013-2015. Volumetric data obtained from cardiac magnetic resonance (CMR), one year prior and about one year after PVR. PS was defined as peak-peak gradient ≥35mmHg by catheterization or peak gradient ≥50mmHg by echocardiography. Student’s t-test and Mann-Whitney test for group comparisons and chi-square and Fisher’s exact test for associations.

Results: 30 patients post-PVR: 15 surgical, 15 transcatheter (1 hybrid). At referral, the SPVR group had only PR; indications for SPVR were: 2+ CMR parameters in 12 patients; 3 had decreased left ventricular ejection fraction (LVEF) and hemodynamic findings and/or abnormal exercise stress test. In the TPVR group, 8 patients had PR, 1 had PS and 6 with PR/PS; indications for TPVR were: 2 CMR parameters in 5 patients, 10 had symptoms and/or hemodynamic findings. At baseline, SPVR group had significantly larger right ventricular (RV) volumes than TPVR group. Biventricular function was not significantly different. Post-PVR, both groups had significantly decreased RV volumes and increased LV diastolic volumes. The SPVR group improved LV cardiac output (CO) (4.9±1.1 to 5.9±0.8 L/min, p=0.03) and biventricular function (RVEF 44.7±5.4% to 53.4±8.8%, p=0.02, LVEF 54.2±5.4% to 60.2±3.8%, p=0.04) whereas the TPVR group had no significant changes.

Conclusions: Most patients were referred for SPVR due to CMR volumetric criteria, whereas for TPVR due to exercise intolerance with mildly abnormal CMR criteria. One year after PVR, both groups had near-normal biventricular volumes and function irrespective of baseline characteristics at referral.
#0057
COMPARISON OF THE LIFETECH CERA AND AMPLATZER VENTRICULAR SEPTAL OCCLUDERS FOR VENTRICULAR SEPTAL DEFECTS CLOSURE IN 147 CHILDREN
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Objective: The Amplatzer and modified double-disc Lifetech Cera ventricular septal defect (VSD) occluders allow the transcatheter closure of the VSDs. The Amplatzer membranous devices are not used anymore because of increased complete atroventricular block risk. Therefore, a comparison of these devices will show us the exact risk of the differences of the devices.

Methods: From May 2009 to July 2016, 147 consecutive patients (mean age 8.5±4.2, range 1.4-26 year) underwent transcatheter closure of VSD. Used devices were Amplatzer membranous in 35 patients (23.8%), Amplatzer muscular in 32 (21.8%), Cera symmetric in 42 (28.4%), Cera muscular in 23 (15.6%), Cera asymmetric in 4 (2.7%) patients. And also 9 patients took Amplatzer ducal occluder I or II and 2 patients Occlutech muscular VSD occluders.

Results: There were no differences in age, sex, defect type and shunt ratio between groups. Membranous defect ratio was 71%. Amplatzer device sizes (7.22±2.11, range 4-16 mm) were bigger than Cera devices (6.22±1.83, range 4-10 mm) (p=0.009). Pacemaker implantation was performed temporary in 3 and permanently in 1 patient at Amplatzer group and none in Cera group. The follow-up period was statistically longer at the Amplatzer group (22.15±16.56 vs 4.76±5.52 months, p<0.001). The other complication and residual flow and success rate were similar at both devices (p>0.05). Membranous Amplatzer devices used in 72.9% of membranous defect but Cera symmetric and asymmetric devices used 94.3% of membranous defects (p<0.001), because of increased block risk of Amplatzer devices.

Conclusion: Although success rate was similar, Cera devices compare favorably with lower complete block risk. And they make more alternative with choosing symmetric, asymmetric and eccentric types.

#0058
INTERVENTIONAL TREATMENT IN ADULTS WITH FONTAN PHYSIOLOGY
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Introduction: The Fontan procedure made increasing survival rate in patients with single ventricle. And early outcomes were reported as good. However, recently the long-term complications have reported by many authors and they need to be regularly followed up with/without catheterization. So, we studied the when and how to do interventional treatment in adults Fontan physiology.

Method: From Jan 2010 to Sep 2015, total 217 patients underwent catheterization in our hospital. Male was 110, and median age of patients was 16.7 years old. Median duration from Fontan procedure was 11.8 years. We reviewed their record retrospectively.

Results: Of them, 118 patients underwent interventional procedure; 15 patients underwent the closure of fenestration in Fontan tract, 58 patients underwent the occlusion of venous collaterals, 34 patients underwent occlusion of arterial collaterals, 31 patients underwent balloon angioplasty, 11 patients underwent stent implantation. There was no intervention- associated mortality. Transient severe headache developed in 1 patient but disappeared in 1 week. It might be due to air emboli or thrombi. Mild hemoptysis occurred in 1 patients but disappeared after 1 day.

Discussion: In spite of good early results of Fontan operation, the long-term complication is not rare and the close follow-up and appropriate intervention is needed. There are many alternative imaging and treatment modalities. However, the well-prepared transcatheter intervention is not risky and good alternative to surgery

#0059
IMMEDIATE AND SHORT TERM OUTCOME POST VSD CLOSURE USING NITOCCLUD PFM COIL – COMPLICATIONS ARE NOT UNCOMMON
Dina Adel, Amr Mansour, Alaa Roushdy, Heba Attya, Azza Elfiky, Maly Elsayed
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Objective: We sought to study the immediate and short term outcome post VSD closure using nitocclud PFM coil to document the safety and efficacy of the procedure.

Patients and Methods: The study included 16 patients with perimembranous subaortic VSD who were scheduled for elective trans catheter VSD closure using nitocclud PFM coil in the period from May 2014 to July 2016. All patients underwent full clinical examination, ECG and full ecochographic study immediately before trans catheter closure as well as 24 hour, 1 month and every 6 month after the procedure. Any intra or post procedural complications and their respective management were recorded.

Results: The mean age of the study subjects was 6.3 ± 3.3 years. The distance between the defect and the aortic valve was an average of 5.4 ± 1.8 mm, and the left ventricular opening averaged 10.6 ± 3.7 mm. Immediate closure of the VSD was acheieved in 25% of the cases this percentage increased to 75% after 1 month. Intravascular hemo-lysis developed 3 days after the procedure in one patient with residual shunt and was successfully managed by PDA ampltazer occluder device implanted in the residual shunt. One patient reported syncope 1 week after discharge and was admitted with complete heart block for which he received transient pacemaker and steroid therapy for 2 weeks after which he regained normal sinus rhythm. The same patient had 2 more attacks of loss of consciousness 2 years after device implantation and complete heart block was documented by ECG and a VVIR pacemaker was eventually implanted for this child. 2 more children had transient self limiting bradycardia and junctional rhythm during the procedure.
Conclusion: VSD closure using nitoclud PFM coil is safe and effective in selected patients. However, we report hemolysis in a patient with residual shunts as well as delayed onset complete heart block in another patient which needs careful follow up and prompt management.

#0060
PRELIMINARY EXPERIENCE WITH THE USE OF THE AMPLATZER VASCULAR PLUG IV (AVP IV) FOR PERCUTANEOUS CLOSURE OF THE SMALL PATENT DUCTUS ARTERIOSUS

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Introduction: Percutaneous occlusion of the small patent ductus arteriosus (PDA) with controlled or uncontrolled release coils can be fraught with residual shunting, mal position and embolization of the device. Although the Amplatzer Vacular Plug IV has been designed to close unwanted vascular structures using low profile catheters, its use for closure of the PDA has not been reported.

Objective: In this preliminary experience, we report the outcomes after the use of this device to close small PDAs in children.

Material and Methods: This is an observational, prospective, longitudinal study with a single cohort of pediatric patients (pts) who underwent percutaneous closure of the small PDA with the AVP IV. Data collection was performed retrospectively through analysis of records. Closure was performed through the retrograde approach using 4 Fr right Judkins diagnostic catheters under general anesthesia. Size of the device was at least twice the size of the minimal diameter of the PDA. Transthoracic echocardiography (TTE) was employed to assess short-term follow up outcomes.

Results: Between 9/15 and 7/16 11 pts underwent closure. Median age and weight was 64.9 (14.1 to 745) months and 20 (7 to 78) kgs, respectively. The mean minimal diameter of the PDA at the pulmonary end was 1.3 ± 0.6 mm and 6.6 ± 2.5 mm at the aortic ampulla mouth. No patient had pulmonary arterial hypertension. The most frequently used device diameter was 4 mm, with a mean of 4.7 ± 1.3 mm. One pt had undergone a previous unsuccessful attempt at closure with a Gianturco coil, which was removed due to malposition inside the aortic ampulla. Technical success and immediate closure was observed in all patients with no complications. Median fluoroscopy time was 3 minutes. During a mean follow up of 6 months, TTE showed closure in all patients with no aortic or left pulmonary artery stenosis.

Conclusion: Percutaneous closure of the small PDA with the AVP IV was feasible and simple from the technical standpoint, safe and effective. Its low profile and possibility of recapture and reposition are the main advantages. Further studies with larger number of pts including infants are warranted.

#0061
PRELIMINARY EXPERIENCE WITH THE HYBRID APPROACH TO MANAGE PATIENTS WITH PULMONARY ATRESIA, VENTRICULAR SEPTAL DEFECTS AND MULTIPLE AORTO-PULMONARY COLLATERALS

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Introduction: Pulmonary atresia with ventricular septal defect and multiple aorto-pulmonary collaterals (PA/VSD MAPCAS) has a broad variety of clinical and anatomic presentation. Surgical repair generally requires a RV to PA conduit with unifocalization of pulmonary blood flow and closure of MAPCAS. Catheter closure of such vessels before surgery may occasionally result in significant desaturation. On the other hand, intra-op ligation of these vessels may be challenging due to their location and friability. Therefore we hypothesized that a hybrid management of patients with this disease could minimize the need of repeat interventions and optimize outcomes.

Objective: To report immediate outcomes after a novel hybrid approach of patients with PA/VSD and MAPCAS.

Material and Methods: This is a prospective, longitudinal, observational study with a single cohort of pediatric patients with PA/VSD and MAPCAS. Data was collected retrospectively with analysis of charts. Patients underwent cardiac catheterization and angio CT to assess pulmonary arteries (PAs) and collaterals before the hybrid approach. Patients were taken to a dedicated hybrid room and a median sternotomy was performed under general anesthesia. Before the initiation of CPB, significant collaterals were closed through an arterial approach via cut-down using a variety of devices, mainly Amplatzer vascular plugs. Surgical repair included BTT shunts or a RV-PA conduit and recruitment of collaterals with Anastomosis to the pulmonary arteries (PAs). The decision to close the VSD was dependent on the size and distribution of the PAs. An exit angio after cessation of CPB was performed if judged appropriate and further interventions were carried out if needed.

Results: Between 01/10 and 8/16, 7 patients with PA/VSD, MAPCAS and small PAs underwent the hybrid approach at a median age and weight of 17.7 (6-102) months and 9.8 (4.2-19.8) kgs, respectively. Two patients had undergone previous palliations including modified BTT shunts and unifocalization procedures. All collaterals were successfully closed before CPB. Six pts had a RV-PA conduit and 1 had a BTT shunt. Mean CPB, anoxic and surgical time was 139.1±37.3, 89.2±23.2 and 494.6±105.2 minutes, respectively. In one patient, PA stenting through the RV-PA conduit was required after exit angio. Pre and post Sats were 77.7±6.6 e 86.6±6.5 % (p< 0.01), respectively. Median ICU, ventilator and vasoactive meds time was 6 days (5-98), 2 days (1-15) and 5 days (0-18), respectively. One patient underwent early post-op catheter closure of a collateral due to significant residual shunting and 2 patients underwent early balloon/stenting angioplasty of the PA. Median in-hospital period was 29 days (7-155). There were no deaths or vascular complications.
Conclusions: The hybrid approach for pts with PA/VSD, MAPCAS and small PAs is a promising new therapeutic modality. Although this approach was feasible and safe in our hands, repeat procedures were still common in the early post-op period. A more aggressive attitude towards intra-op rehabilitation of the PAs after CPB may be required as experience evolves.

#0062
CATHETER CLOSURE OF PATENT DUCTUS ARTERIOSUS IN ADULT PATIENTS USING AN OUTPATIENT PROTOCOL
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Introduction: Transcatheter closure is the treatment of choice for the majority of patients with a patent ductus arteriosus (PDA). However, the standard technique of this procedure uses an arterial access and requires immobilization of the patients for 24 hours and may be associated with arterial complications. The aim of this study was to report experience with catheter closure of PDA in 68 consecutive adult patients the Cocoon PDA occluder on outpatient basis using an exclusive venous approach.

Methods: The age of the patients ranged from 16-72 (median 36 years) and the weight from 52-74 Kg (median 64 Kg). The anatomy and size of PDA were defined by transvenous retrograde aortography using a Pigtail or a Berman catheter. The PDA occluder was implanted through an 8-9 F delivery sheath (DS). The procedure was guided using hand injections of contrast media through the DS and 2D and color Doppler echocardiography from suprasternal and parasternal longitudinal and short axis, respectively.

Results: The PDA occluders were permanently implanted in 65/68 of the 68 patients. The mean PDA diameter (at the pulmonary end) was 3.8±0.9 mm (range 1.2 to 9.8 mm). The mean device diameter was 6 ± 3 mm (range 4 to 12 mm). Complete echocardiographic closure of the ductus at 1-month follow-up was observed in all 65 patients (100%). Eight minor groin venous hematomas were the only complications of the procedure. The hospital stay of the patients ranged from 6-8 hours.

Conclusions: Exclusive transvenous PDA occlusion using combined angiographic and echocardiographic guidance is an effective and safe method that prevents the arterial complications of the standard approach. In addition, in adult patients, this technique, may be used on outpatient basis resulting in reduce hospital cost.

#0063
PERSISTENT PLEURAL EFFUSION, PULMONARY STENOSIS, FAILED FONTAN CIRCULATION. “HEADACHE” SOLVED IN THE CATH LAB.
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A 6 year old girl with a history of double discordance, narrow bulboventricular foramen and hypoplastic right ventricle, had been surgically palliated at one year to ensure systemic flow to the aorta, through Aortopulmonary window with main pulmonary artery banding to reduce the flow and pressure to the lungs and guarantee the systemic output. At 4 years catheterization and later Glenn performed, it was decided to keep intentionally the anterograde flow through the pulmonary artery. At 6 years was completed the cavopulmonary derivation, Cath hemodynamic parameters: (Main Pulmonary Artery: 13/10, mean:10mmHg, Left Ventricle: 80/8 mmhg, Index Pulmonary Vascular Resistance 1.2 Wood Units, pulmonary branches normal Z score). She underwent extracardiac Fontan with fenestration, at the same time the main pulmonary artery was ligated. The immediate outcome was satisfactory. Almost to completed the second week post-surgical the right pleural drainage increase. Echocardiogram reveal anterograde flow through the pulmonary artery and stenosis at the confluence of the pulmonary artery and proximal third of the left branch. She stayed hospitalized for 10 weeks, high right lung drainage persists even though medical therapy was optimized and local procedures were conducted. Cath findings were: the proximal third of the left branch was 6 mm (Z-4.4) and the distal branch was 10.6 mm (Z-0.17).

We realize that both superior and inferior vena cava had selective flow to the right pulmonary branch. Fenestration was not evident. Hemodynamic findings: Superior and inferior vena cava mean pressure 20mmHg, and distal prebranching left artery 15mmHg. Ediasole ventricular pressure was 9 mmHg. PROCEDURE: A test occlusion in the pulmonary artery using a noncompliant balloon (simulating delivered covered stent) and simultaneously angiography in ascending aorta to confirm the occlusion of the anterograde flow through the aorto-pulmonary window was carried out, and we assess the free drainage of the superior and inferior vena cava. By using a 12Fr Mullins long sheath we implanted a CP Stent™ covered 10 Zigs of 39 mm, mounted on 12x40 mm POWERFLEX®Pro balloon, and later stent redilation to 14 mm with MAXI LD™ balloon, without complication. Post angiography result was evaluated, finding homogeneous flow in both pulmonary branches and the Fontan pressure was the same throughout the system, the accessory pulmonary flow was totally blocked. Three days later, the pleural drainage was minimum and she was discharge 10 days after the intervention.

Conclusion: There are numerous causes that can complicate a patient with Fontan circulation, persistent pleural effusion is one of the most common, it’s imperative to perform catheterization early in patients with failed Fontan. Nowadays we should seek for the procedure that brings the better outcome and with less morbidity and mortality to our patients such as the case we discussed, in which discharge was possible 1 week later, after a long hospital stay.

#0064
PERCUTANEOUS AORTIC PARAVALVULAR LEAK CLOSURE USING AMPLATZER VASCULAR PLUG III
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Paravalvar leak (PVL) is a relatively common complication of valve replacement surgery. Although most PVLs are small and
Asymptomatic, 2% to 5% are clinically relevant and associated with major complications, such as heart failure, hemolytic anemia, arrhythmias, and infective endocarditis. Surgical reoperation is effective however associated with morbidity and mortality especially in high risk patients. To date percutaneous PVLs closure has proved high efficacy, with different device or occluders, the most used is amplatzor vascular plug III. We report a 40 year male with aortic bivalve, and severe aortic regurgitation who underwent 2 years ago aortic valve replacement with magna ease bioprosthetic aortic valve, after surgery developed heart failure, arrhythmias. Echocardiography showed severe aortic paravalvular leak, Under general anesthesia, transesophageal echocardiography guidance (TEE) trough right femoral artery cardiac cath was performed to close the leak, angiography and TEE evidenced a leak on left coronary sinus, measures long axis 12 mm, short axis 4 mm. AL 2 catheter and hydrophilic wire was used to cross the leak. With exchanged wire in left ventricle an amplatzor vascular plug III 14x3 mm was delivered, placed in the leak and release, control angiography and TEE showed good position and minimal residual shunt, the patient was discharged next day. On follow up 8 months the patient refers clinical improvement, is not receiving anhyrarrhythmic or diuretic. Echocardiography normal left ventricular function, mild LV dilation, and mild residual PVL. We report a successful Amplatzor vascular plug III aortic PVL closure with good clinical improvement, and mild peri device residual leak.

Key words: Amplatzor vascular plug III • aortic paravalvular leak • percutaneous

#0065
CLOSURE OF LARGE AND LONG TUBULAR PATENT DUCTUS ARTERIOSUS IN INFANTILE AGE GROUP USING THE AMPLATZER® VASCULAR PLUG II
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Objective: To evaluate the safety and efficacy of the Amplatzor Vascular Plug II (AVPII) in closing large and long tubular Patent ductus arteriosus (PDA) in infants and young children.

Background: Large and long tubular PDAs in young infants (Figure 1a & 2a) are not amenable for closure with the conventional duct occluders and necessitate surgical closure. Previous case series and reports have described use of AVPII for closure of such PDAs in premies or for smaller sized ducts in younger infants. We describe the use of AVPII for closure of such large and long PDAs in young infants (Fig 1b & 2b).

Methods: Data of patients undergoing transcatheter PDA closure between May 2015 & August 2016 were retrospectively reviewed. Transthoracic echocardiography (TTE) was done in all cases to confirm the diagnosis as well as to characterize the PDA in terms of size at pulmonary (PA) and aortic end, shape, length and gradient across the PDA and size of aortic isthmus and left pulmonary artery. Procedural steps, complications and results were recorded.

Results: Ten patients (5 females) underwent transcatheter PDA closure. The mean age was 7.3 ± 4.7 months and mean weight was 5.4 ± 1.5 kg. All patients had shortness of breath and 9 of 10 had evidence of failure to thrive. The PDA measured a mean of 5.1 ± 1.0 mm at the PA end with an average length of 10.4 ± 2.8 mm. All patients had evidence of large shunt with the mean aortic pulse pressure being 55.4 ± 8.9 mm Hg. The mean difference between the aortic and pulmonary systolic pressure heads was 16.2 ± 9.6 mm Hg suggestive of minimally restricted ducts. The AVPII size ranged from 6 to 10 mm. The mean device to PDA size ratio was 141 ± 9.5%. The device was deployed from the venous side in all cases and deployed completely within the duct. Successful occlusion of the PDA was achieved in 8 out of 10 cases (80%) with no evidence of aortic or left pulmonary artery obstruction. In one case the device embolized after release and the child was shifted for surgical device retrieval and PDA closure. In another case, the device was felt to be unstable after deployment and hence was not released. The child was sent for surgical PDA ligation electively. The average device length to PDA length ratio in both the cases was 102.5% compared to an average of 59.1% in those where the procedure was successful. At a mean follow up of 10.1 months (range 1-15 months), all patients were doing well with no evidence of any residual shunt on TTE. The LPA and the aortic isthmic flows were laminar with no evidence of obstruction.

Conclusion: We report successful use of AVPII for closure of large, long and tubular PDAs with significant pulmonary arterial hypertension in young infants. The device tends to retain its position after intraductal deployment even in the absence of a retention disc by virtue of ductal walls offering the support to stabilize the device. Inadequate ductal length can result in the device getting unstable and may lead to embolization.

#0066
LEAFLET MORPHOLOGY VARIATION OF THE MELODY TRANSCATHETER PULMONARY VALVE: EFFECT ON PERFORMANCE AND OUTCOME
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Background: The Melody® Transcatheter Pulmonary Valve (TPV) is constructed using a harvested bovine jugular venous valve, which is rigorously tested for competency prior to commercial release. Natural anatomic leaflet variations are seen in the TPV when viewed en face. These valve morphologies have not been described, nor evaluated in the TPV.

Methods: A TPV morphology classification system was devised after reviewing 1/3 of the available photographic images of implanted TPVs at our institution. All images were blindly reviewed by implanters and TPVs were classified based on the consensus of the reviewers. The TPV outcomes (echocardiographic function and complications) were compared based on leaflet morphology. Complications included endocarditis, stent fracture, TPV replacement (surgical or transcatheter), and patient death.

Results: A total of 62 TPVs were categorized into the following leaflet morphology types: A -symmetric trileaflet (47%); B-asymmetric trileaflet with a single small leaflet (32%); C-asymmetric trileaflet with a single large leaflet (16%); D-rudimentary leaflet with near bicuspid
appearance (5%). Seven TPVs were excluded from analysis as they were not implanted in the pulmonary valve position. Only one TPV (Type C) had >mild regurgitation on the first post-procedural echocardiogram. Over a median follow-up of 1.9 years (range 0.02-7.8 years), 9 TPVs developed complications with 7 requiring re-intervention. The majority of complications were not attributed to the valve morphology including right ventricular outflow tract obstruction (RVOTO)/valve stent fracture (n=4, 1 with pre-stented), and ventricular arrhythmia (n=1). Endocarditis was diagnosed in 3 patients (2- Type B and 1- Type A), one resulting in death. Two TPVs (2- Type A) had >mild regurgitation over the follow-up period and both were secondary to stenosis associated with a complication (endocarditis and RVOTO). There was one patient death of unclear etiology (Type B). There was no significant difference in outcomes based on TPV leaflet morphology type.

Conclusions: The Melody® TPV can be classified into one of four categories based on leaflet morphology. In this cohort, there was a low incidence of complications which were not associated with leaflet morphology. Implanters should be aware of the variation in TPV morphology, and documentation may lead to better understanding of associated outcomes.

#0067
NOVEL APPROACH TO TRANSVENOUS PACING IN THE FONTAN CIRCULATION
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Introduction: The need for pacing in the Fontan circulation increases with age. Although epicardial pacing is most common, transvenous pacing is required in some.

Objective: To describe a novel approach to transvenous pacing in a patient with non-fenestrated Fontan circulation.

Methods: An 18 year-old male with hypoplastic left heart syndrome was palliated with an intracardiac lateral tunnel. His surgical fenestration and intracardiac baffle leaks were previously occluded. Due to progressive high grade AV block, an epicardial system was placed, upgraded to a cardiac resynchronization system, and revised three times for lead failure. During the last revision, his sixth sternotomy, dense adhesions limited surgical access and adequate pacing sites. He presented with need for pacemaker revision secondary to increasing impedance thresholds; a surgical approach was deemed prohibitive. A novel approach to gain access to the systemic circulation for ventricular pacing was proposed. A left-sided pre-pectoral pocket was created. The left subclavian vein was accessed using Seldinger technique. An 8 Fr Bard Channel steerable sheath was advanced to the floor of the pulmonary artery. The system was exchanged over a guidewire and the tract was balloon dilated allowing placement of a delivery sheath into the systemic ventricle and a bipolar active fixation lead was placed and an atrial lead was placed on the free wall of the lateral tunnel.

Results: The patient made an uneventful recovery and has excellent pacing thresholds. He is anticoagulated and in the short term (3 months) has not suffered a thromboembolic event. There is no significant systemic AV valve regurgitation.

Conclusion: This innovative method provides an option for transvenous pacing in patients with non-fenestrated intracardiac or extracardiac Fontan connections when epicardial leads are not feasible. The long term risk of thromboembolism and AV valve functioning will need to be evaluated.

#0068
NOVEL TECHNIQUE IN RIGHT VENTRICULAR OUTFLOW STENTING IN AN EXTREMELY SMALL PREMATURE INFANT WITH TETRALOGY OF FALLOT
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Right ventricular outflow tract stenting has had significant advances since the first stents were placed into surgically placed right ventricle to pulmonary artery conduits in the early 1990s. Catheter and stent technology have made it increasingly possible to treat smaller infants, more recently with echocardiography guidance. There have been multiple series reports of successful palliations using these cardiac procedures in larger patients, using easily available equipment. Smaller patients present an additional challenge to an already complex situation.

We discuss a case of a 1.3kg premature infant who underwent complex right ventricular outflow tract stenting as an emergent procedure by employing a novel technique using neurovascular catheters, to aid wire exchanges.

#0069
PREDICTORS OF PROCEDURE TIME PROLONGATION DURING PERCUTANEOUS TRANSCATHETER CLOSURE OF ATRIAL SEPTAL DEFECT; A RETROSPECTIVE STUDY
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Mohammed Omar Galal⁴
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Background: Percutaneous transcatheter closure (PTCC) of atrial septal defect (ASD) may convert to a long procedure. We aimed to identify predictors of prolonged procedure.

Methods: Under transesophageal echocardiography and fluoroscopy guidance, 81 children with ASD underwent PTCC. Retrospectively,
Results: The procedure was prolonged in 25 patients. By mono-
variate analysis, the significant predictors for prolonged procedure
were smaller and younger patients, larger ASD, smaller left atrial
(LA) dimensions and device waist ratios to patient’s length, and
LA dimensions. By multivariate analysis, the significant predic-
tors were deficient septal rim toward superior vena cava (SVC) and
device waist diameter in relation to patient’s length (best cut-off:
< 12 mm and > 0.13, respectively). Presence of Chiari network
was not statistically significant, but it prolonged procedure duration.
In three cases (3.7%) the device embolized; all had short SVC rim.

Conclusions: A short septal rim toward superior vena cava and large
device waist size in relation to patient size and/or LA dimensions may
predict prolonged procedure during PTCC of ASD. Chiari network
may also be a risk factor. Short septum rim toward superior vena cava
may be a risk factor for embolization.

#0070
PERCUTANEOUS VSD CLOSURE UNDER 1 YEAR OF AGE:
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Suleyman Sunkak, Kazim Uzum
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Background and aim: Untreated large Ventricular septal Defects(VSD)
are important reason of congestive heart failure in early infancy.
This population usually fails to grow and surgical closure is challeng-
ing because of congestion in their lungs prone to respiratory infec-
tion and their bad nutritional status. We therefore planned to close
VSD of such patient group under 1 year of age percutenously and
wanted to share our experience.

Material & Method: We have performed VSD closure of 7 patients
under 1 year of age between the dates September 2012- May 2016 in
Erciyes University Pediatric Cardiology Department.

Table 1 (#0070): Demographic and angiographic Data of Patients

<table>
<thead>
<tr>
<th>Patient</th>
<th>Age(months)</th>
<th>Weight(kg)</th>
<th>VSD Type</th>
<th>VSD diameter(mm)</th>
<th>Device Type</th>
<th>Device Size</th>
<th>Fluoroscopy time(min)</th>
<th>Radiation Dosage</th>
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<tbody>
<tr>
<td>1</td>
<td>9</td>
<td>7</td>
<td>muscular</td>
<td>3.5</td>
<td>ADO-II</td>
<td>5x6</td>
<td>32</td>
<td>1747</td>
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<tr>
<td>2</td>
<td>4</td>
<td>5.3</td>
<td>Perimembranous</td>
<td>3</td>
<td>ADO-II</td>
<td>4x4</td>
<td>22.6</td>
<td>1027</td>
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<tr>
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<td>9</td>
<td>8</td>
<td>Perimembranous</td>
<td>4</td>
<td>ADO-II</td>
<td>6x4</td>
<td>15</td>
<td>495.7</td>
</tr>
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<td>ADO-II</td>
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<td>ADO-II</td>
<td>3x4</td>
<td>14</td>
<td>1002</td>
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</tbody>
</table>

Results: Age of patients ranged between 4 months-12 months.
Weight of the patients during the procedure was between 5.3-9
kg. Mean VSD diameter was 3.19±0.47mm. One of defects was
muscular, rest of them were perimembranous. All defects were
closed with Amplatzer Ductal occluder II(ADO-II). Mean fluoros-
copy duration and total radiation dosage were 78.5±94.6min,
2069±1395cGy/min respectively. We did not faced with any
major complication except in one patient: complete AV block was
seen one month after the procedure. Pacemaker was implanted.
No aortic regurgitation was seen in patients after device
implantation.

Conclusion: The procedure of VSD closure, whether it is surgical or
percutaneous, is very risky. The risks were higher when the children
were smaller than 1 year of age and low body weight. Percutaneous
VSD closure may be an alternative to surgery in early infancy that
carry the similar risks but less invasive.

#0071
PERCUTANEOUS PDA CLOSURE IN EXTREMELY LOW
BIRTH WEIGHT BABIES
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Ayse Ulgey, Kazim Uzum
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Aim: Patent Ductus Arteriosus(PDA) is an important cause of morbid-
ity and mortality in preterms. As birthweight decrease, risks increase.
Main aim of our study is to emphasize the effectiveness and safety of
percutaneous PDA closure even in extremely low birth infants (less
than 1000gr)

Material Method: In our center between the dates June2014-
December2015, PDA of eight patients less than1000gr were closed
percutaneously. To our knowledge this study includes the largest
cohort of infants less than 1000g in the literature, whose PDA were
closed percutaneously.

Results: Symptomatic patients, less than1000gr having PDA were
included in the study. All have3times medical therapy for PDA closure
but it didnotwork. PDA was decided to be contributor of this medical
state of them. The mean patient age 16±5.9days. The mean weight of
patients was 923±75.9gr. Mean gestational age was 27.2±1.28weeks.

Conclusions: A short septal rim toward superior vena cava and large
device waist size in relation to patient size and/or LA dimensions may
predict prolonged procedure during PTCC of ASD. Chiari network
may also be a risk factor. Short septum rim toward superior vena cava
may be a risk factor for embolization.
Mean PDA diameter was 2.48±0.5mm. Mean Qp/Qs was 1.7±0.2. Morphology of PDA: 50% of them were conical, 30% of them were tubular. In all patients ADIOI-AS device were used for PDA closure (Table 1). Steps of percutaneous PDA closure procedure was shown by Figure 1. In all patients, we have done closure by venous route. We did not ever used arterial route in 4 patients. There were no major complications reported. Left pulmonary arterial stenosis was detected in 2 patients which were all resolved in 6 months duration.

Conclusion: Interventional catheterization procedures are more commonly used, in the recent years. The advantages of percutaneous PDA closure include a high success rate, shorter length of hospital stay, reduced blood loss, low morbidity rate, and no traumatic scars. Since the length of hospital stay decreases with catheterization, it is much more cost-effective than surgery. We want to emphasize that in experienced centers percutaneous closure of PDA can be an alternative to surgery even in the extremely low birth weight babies.

#0072
INTENTIONAL LACERATION OF THE ANTERIOR MITRAL LEAFLET TO PREVENT LEFT VENTRICULAR OUTFLOW TRACT OBSTRUCTION (LAMPOON) DURING TRANSCATHETER MITRAL VALVE IMPLANTATION: PRE-ClinICAL FINDINGS
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Background: Left ventricular outflow tract (LVOT) obstruction is a life-threatening complication of transcatheter mitral valve implantation (TMVR), caused by septal displacement of the anterior mitral leaflet (AML). We propose a novel transcatheter resection of the AML.

Methods: LAMPOON was developed in vivo in swine, guided by biplane X-ray fluoroscopy and intracardiac echocardiography. Two transfemoral coronary guiding catheters were advanced retrograde through the aortic valve, either side of the AML. The ‘LVOT catheter’ directed an electrified guidewire across the center and base of the AML towards a snare directed by the ‘left atrial catheter’. The snared externalized guidewire loop was electrified, lacerating the AML along the centerline from base to tip.

Results: The AML was successfully lacerated in all (n=8) swine. Benchtop analysis demonstrated an increase in neo-LVOT with some residual obstruction (65% ± 10% vs. 31 ± 18% patency, p<0.01). Mean blood pressure fell (54 ± 6 to 30 ± 4 mm Hg, p<0.01) due to acute mitral regurgitation from LAMPOON, but remained steady until planned euthanasia. There was no collateral tissue injury on necropsy.

Conclusions: Using simple catheter electrosurgery techniques, we mimicked surgical chord-sparing AML resection prior to TMVR. We recommend pre-positioning the transcatheter heart valve to minimize time between laceration and TMVR, though laceration may be better tolerated in patients with chronic mitral valve disease. Applied cautiously, this technique may enable TMVR in high risk surgical patients with prohibitive risk of LVOT obstruction.

#0073
POST-IMPLANTATION MODIFICATION OF ENDOVASCULAR STENTS IN THE TREATMENT OF COMPLEX CONGENITAL AND ACQUIRED VASCULAR STENOSES
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Background: Optimal treatment of complex congenital and acquired vascular stenoses may benefit from post-implantation modification of standard endovascular stents. Utilization of open-cell design stents allows for side-hole creation with dilation of stent struts to minimize impact on adjacent vessels and/or treat adjacent stenotic lesions. We sought to describe our experience with this technique.

Methods: Single-center retrospective review at a large congenital heart center from 2011 – 2016. Cases with endovascular stent placement followed by side-hole creation with or without additional stent implantation, or with side-hole modification of an existing implant, were included. Demographic and procedural details were collected.

Results: Twenty catheterization procedures in 19 patients met inclusion criteria. Cases were performed in 55% female patients with a median age of 12.8 (IQR 4.2, 25.4) years and a mean weight of 45.1 ± 33.3 kg. Indications for stent angioplasty included: complex RVOT and branch pulmonary artery (PA) stenosis (45%), lobar PA stenosis (30%), aortic coarctation (15%), atrio-pulmonary Fontan connection stenosis (10%), and systemic venous stenosis (10%). During 18 procedures a new stent was placed, while an existing implant was modified in 2 procedures. Three types of open-cell stents were modified following implantation: ev3 IntraStent® LD Mega (45%), ev3 IntraStent® LD Max (35%) and Cook Formula 418 (30%). A lobar (45%) or branch (30%) PA were the most commonly jailed vessels, followed by left subclavian artery (15%), right internal jugular vein (5%) and hepatic vein (5%). Indications for side-hole dilation included: relief of side-branch stenosis caused by primary stent implant (25%), avoidance of stent material (struts) crossing an important side-branch (50%) and relief of pre-existing side-branch vascular stenosis (25%). Side strut angioplasty was performed to a variety of diameters (4 mm – 24 mm) using high pressure or ultra-high pressure angioplasty balloons. In 4 cases, a second stent was placed through the newly generated side-hole. Median procedure time was 211.5 (123.5, 272.5) min and mean fluoroscopy time was 56.2 ± 23.8 min. There were no local vascular injuries related to post-implant stent modification but 1 patient did have a self-limited access site complication.

Conclusion: Post-implant modification of endovascular stents via side strut dilation is safe, feasible and may improve outcomes in the definitive treatment of complex vascular lesions.

#0074
PERCUTANEOUS CLOSURE OF POSTMYOCARDIAL INFARCTION VENTRICULAR SEPTAL DEFECTS
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Background: Interventricular septum rupture following acute myocardial infarction (MI) is a rare (≈0.2%) but serious and potentially life threatening complication. Current AHA/ACC guidelines recommend urgent surgical repair even in the absence of hemodynamic compromise. However, surgical mortality rates remain high (reported between 20 to 80%) particularly in patients with cardiogenic shock. Percutaneous intervention is an attractive, less invasive alternative yet this approach remains to be performed only in few selected centers around the world.

Methods: Single centre, retrospective study of 17 consecutive patients diagnosed with postmyocardial infarction ventricular septal defect (pMIVSD) that underwent percutaneous device closure between July 2003 (index procedure) and July 2016. Patients were further divided into those with (n=8) (Group A) and without cardiogenic shock (n=9) (Group B) at the time of intervention. The latter was defined as those requiring inotropic support or intra-aortic balloon counterpulsation to sustain cardiac output.

Results: Nine (47%) patients were female. Median age was 65 years (range 51-75 y). Median time in days between MI and pMIVSD closure was 8 days (range 1-25) for group A and 11 days (range 3-91) for Group B. Systolic pulmonary artery pressure/systemic pressure ratio was 0.57 (range 0.38-0.7) and Qp:Qs 1:3.1 (1.72-3.6) for patients in Group A and 0.6(range 0.33-0.89) and Qp:Qs 3:1 (1.58-4.39) for patients in group B (NS). A variety of devices were used to occlude the defect: muscular VSD Amplatzer n=6(35%), postinfarct VSD Amplatzer n=5(30%), ASO n=4(24%), other (AVP II and PFO) n=2(11%). Device embolization occurred in only one patient (device was snared and retrieved) and a second occluder used. In group A, significant residual shunt was found in two cases; hence they were brought back to the lab to deploy a second occluder. No reintervention was required for patients in group B, mild to moderate residual shunt was present in 6 out of 9 (66%) patients. Early (<30 day) mortality was 0% in Group B and 62% (5 out of 8 patients) in Group A.

Conclusion: Transcatheter device closure of pMIVSD can be considered a reasonably safe and effective procedure in selected patients. Our series is consistent with previous experience reported in other series regarding high mortality in patients with cardiogenic shock despite intervention.

#0075
THE IMPACT OF TRANSCATHETER PULMONARY VALVE IMPLANTATION ON THE MANAGEMENT OF POSTOPERATIVE RIGHT VENTRICULAR OUTFLOW TRACT DYSFUNCTION
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Pulmonary valve implantation (PVI) for pts with CHD has undergone revolutionary changes in recent years with the advent of transcatheter therapies. Surgical and transcatheter procedure characteristics and valve function were investigated in 172 consecutive pts referred for PVI (median weight 56.7kg [12.3-161.3]; median age 18.6 years [4.7-65.5]) from 2007 to 2016. Diagnoses included tetralogy of Fallot (126), PA/PS (IVS) (21), and other (25). Indication for PVI was PI in 107, PS in 31 and PS/PI in 34. The number of PVI procedures increased from 9 to 27 per year with the availability of transcatheter pulmonary valve implantation (TcPVI). Prior to the availability of TcPVI, 34 pts were referred for surgery. Of the next 138 pts, 101 were referred for TcPVI and 37 went directly to surgery. TcPVI was attempted and successful in 85/101. The remaining 16 were referred for surgical pulmonary valve implantation (sPVI) without any attempt to implant a percutaneous valve due to RVOT size (11), coronary or aortic compression (4) and conduit tear pre-PARCS (1). 8/11 pts referred for surgery due to RVOT size would have been candidates for the 29mm Sapien valve if it had been available. TcPVI was successful in 27/27 pts with a bioprosthetic pulmonary valve (BPV), 24/27 with homografts and 34/47 with transannular patch repair (TAP). A Melody valve was implanted in all pts with a BPV or homograft and in 22/34 TAP pts. The other 12 pts received a Sapien valve. In total, 53 pts were referred for surgery in the TcPVI era. The RVOT was too large for TcPVI (44), too small (4), coronary or aortic compression (4) and conduit tear (1); 2 declined surgery. Comparing TcPVI to sPVI in the same time frame, freedom from > mild PI was 84/85 (TcPVI) and 36/51(sPVI) p<.001; 4/51 sPVI underwent subsequent Melody valve implantation for PI. Obstruction developed in 1 pt in each group p=ns. A second Melody valve was implanted in the 1 TcPVI pt with obstruction. SBE occurred in 1/85 (TcPVI) vs 2/51 (sPVI).

Since the introduction of TcPVI, referrals for PVI have tripled and 63% have had a percutaneous approach. Pts treated with TcPVI have had better long-term valve function in comparison to sPVI over the same time period. Given the superiority of TcPVI and the ability to avoid repeat open heart surgery, earlier intervention to reduce the incidence of RV dysfunction and development of symptoms may be warranted.

#0076
THROMBOPROPHYLAXIS STRATEGIES FOR CHILDREN WITH SINGLE VENTRICLE CIRCULATIONS (SUPERIOR OR TOTAL CAVO-PULMONARY CONNECTIONS) AFTER STENT IMPLANTATION
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Background: The most effective thromboprophylaxis strategy after stent implantation in the superior or total cavopulmonary connection (SCPC or TCPC) is unclear.

Methods: We reviewed our cardiac database over a 13-year period ending in 2015. Patients were divided into those who received aspirin (ASA) alone versus advanced anti-coagulation (AA), including warfarin, enoxaparin, heparin, or clopidogrel. Primary endpoint was presence of thrombus on advanced imaging, such as cardiac catheterization, CT-angiogram or MRI. Other endpoints included presence of thrombus on echocardiogram and bleeding complications.
Introduction: Patients with surgically corrected tetralogy of Fallot (TF) often develop severe pulmonary regurgitation (PR) with chronic right ventricular volume overload, leading to adverse outcomes. We created an ovine survival model simulating the pathophysiology of TF to study the effects of right ventricular remodeling due to stenosis and regurgitation.

Methods: In lambs, a pulmonary valve stenosis (PS) was created by placing a PTFE strip around the pulmonary artery through a right thoracotomy (Figure A). Four months later a bare metal stent was anchored across the pulmonary valve in the PTFE strip (Figure B) thereby relieving the stenosis and creating pulmonary valve insufficiency (Figure C). Melody valve implantations into this bare metal stent at different time intervals are ongoing (Figure D). Follow up by means of MRI was performed to assess remodeling and reversed remodeling of the RV.

Results: All animals survived the initial surgical phase (n=9). Two animals died during bare metal stent implantation (ventricular fibrillation n=1; PA rupture by balloon dilation n=1). MRI showed signs of RV hypertrophy prior to relief of stenosis compared to healthy controls. Total RV cardiac output (CO) was 2.2±0.7 L/min after PS, 5.0 ±0.8 L/min immediately after bare stent implantation and 3.5±0.1 L/min SD after 5 months of PR. Animals had an important PR 5 months after bare metal stent implantation (32±2.3%). The LV-RV EDV ratio was 0.7±0.1. This was significantly smaller compared to healthy controls (p<0.05)(n=3).

Conclusion: The creation of an ovine survival TF model with initial pulmonary valve stenosis and secondary regurgitation (mimicking the effect of surgical repair) is feasible. All hallmarks of the TF physiology (ventricular hypertrophy after stenosis – ventricular dilatation due to PR) were realized. This model forms the ideal basis to study the timing of pulmonary valve replacement in TF.

#0078
10 YEARS OF EXPERIENCE WITH THE MELODY™ VALVED STENT FOR PPVR

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Results: A total of 58 patients with single ventricle had 72 stents implanted in the LPA (n=41, 57%), RPA (n=7, 10%), Fontan baffle (n=19, 26%), SVC (n=4, 6%) or innominate vein (n=1, 1%). Of those, 14 stents (19%) were implanted post SCPC and 58 (81%) were after TCPC. The most common indication for stenting was vessel/conduit stenosis (n=61, 85%), followed by intraluminal thrombosis (n=11, 15%). Interval between SCPC/TCPC surgery to stenting was x (IQR y – z) years. AA was prescribed for 32 (44%) patients, while 40 (56%) were treated with only ASA. There were higher rates of pre-stent thrombosis in patients who were treated with AA when compared to ASA (28% vs 5%, p=0.009). Patients on AA had higher rates of post-operative stenting (within 30 days of surgery) (47% vs 15%, p=0.003) and had more severe stenosis (58% vs 43%, p=0.001) when compared to ASA.

Median patient follow up was 1.1 (IQR 0.5 – 2.6) years. Advanced imaging was obtained on 44 patients (61%), with no significant difference between the ASA and AA group (58% vs 64%, p=0.629). Median interval between stent implantation to advanced imaging was 1.2 (IQR 0.6 – 2.1) years. Follow up echocardiogram was available on 71 (99%) of patient. Median interval between stent implantation to echocardiogram was x (IQR y – z) years. Only 2 patients (3%) were found to have intra-stent thrombus at 1 and 3 days post stenting despite being on AA (therapeutic heparin), detected initially on echocardiogram and confirmed by angiography in the cath lab. Both were stented within 3 days post-SCPC surgery due to occlusive LPA thrombus. They both underwent stent re-dilations in the cath lab, and one required surgical thrombectomy, stent removal and LPA arterioplasty. There were 8 significant bleeding complications in the AA group and none for ASA (p=0.005).

Conclusions: In our limited cohort, we found no difference in the rate of intra-stent thrombosis between ASA and AA treatment. ASA alone may be sufficient therapy for most SCPC and TCPC patients undergoing stent implantation, while pre-existing thrombus may warrant AA.

SUCCESSFUL CREATION OF AN OVINE TETRALOGY OF FALLOT MODEL

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A/ PTFE strip supra-annular and marker infundibular.
B/ MRI showing bare metal stent across the native pulmonary valve.
C/ MRI phase shift axis showing massive PR.
D/ Deployment of the Melody™ valve in bare stent.

Figure 1 (#0077). Panel A. PTFE strip supra-annular and marker infundibular. Panel B. MRI showing bare metal stent across the native pulmonary valve. Panel C. MRI phase shift axis showing massive PR. Panel D. Deployment of the Melody™ valve in bare stent.
Introduction: Since 10 years the Melody™ valved stent is used for percutaneous pulmonary valve replacement (PPVR) with over 10,000 implants worldwide. It has been proven to be a safe and effective intervention. The main risk factors of the valve compared to surgical alternatives are stent fractures with subsequent valve dysfunction and endocarditis. Most biological valves have limited long term preserved function requiring redo interventions.

Aim: evaluation of the valve function and its risks from a prospective ongoing registry

Patients and methods: All data are collected from the Belgian registry which started in 2006. Follow-up data are prospectively and ongoing collected. 97% are pre-stented in this cohort. The analyzed data: leaflet function by means of echocardiography, re-interventions, mortality, endocarditis (Duke's criteria) and stent fractures by means of X-ray. Data presented is single center experience.

Results: 159 valves have been implanted 2006-2016. The indication was stenosis 44.0%, regurgitation 33.3% and mixed 22.6%; male/female ratio was 2:1, the mean age at implant was 19.6 y (3.5 – 81.6). Original conduit diameter was median 22 mm (10-26 mm).

1. Overall survival was 98 % at 10 years, there were no procedural or valve related deaths.

2. Endocarditis occurred in 9.4% (15/159) with a freedom from endocarditis of 78% at 10 y. The maximal annual incidence of endocarditis was 3.3% in 2015. Mean time of occurrence of endocarditis was 2.4 y (0.7 – 8.9) after implant. Mean age at endocarditis was 18.2 y (8.1 – 45.6), male 87%.

3. Chest X-ray was performed in 87 % of the patients showing a stent fracture in 8 % median 2.9 y (0.4 – 6.9) after implant. None of the fractures led to hemodynamic problems and no re-intervention was necessary.

4. Re-intervention overall 11 %: explantation of valve in 4.4% (endocarditis n= 6, residual subvalvular stenosis n=1); balloon dilation due to somatic growth in 5 % and 1.9 % received second PPVR (post endocarditis).

5. Valve function: a) indication PS : mean PS 65 mmHg (SD 17) at implant drops to 23 mmHg (SD 11) post implant and is 37.5 mmHg (SD 17) 10 y later. B) Indication PR: median 4/4 at implant drops to 0/4 post and is 2/4 at 10 y follow-up.

Discussion: Ten year follow-up of the Melody™ valved stent shows good long term preservation of the valve function with current no replacement for intrinsic valve dysfunction. No hemodynamic important stent fractures occur since the pre-stenting era. Acceptable risk of endocarditis remains in male young adult patients.

#0079
PERCUTANEOUS DOUBLE VALVE IMPLANTATION IN A GROWN-UP CONGENITAL HEART WITH DEGENERATED HOMOGRAFT IN BOTH AORTIC AND PULMONIC POSITION.
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This is the case of a 43 yr-old male with a truncus arteriosus who underwent three previous cardiac surgeries: - 1972: pulmonary banding; - 1979: prosthetic valve tube between right ventricle and pulmonary artery + aortic valve prosthesis ( Bjork) + VSD closure; - 1991: homograft EHB 22 mm in pulmonic position, homograft EHB 22 mm in aortic position replacing the Bjork prosthesis. He was admitted in April 2016 in the intensive care unit for a severe heart failure. Echocardiography showed a severe aortic regurgitation, a pulmonic valve stenosis, enlarged and poor function of right and left ventricles. Diagnostic cardiac catheterization showed a heavy calcified and stenotic (minimum diameter : 10 mm; RV-PA gradient : 44 mmHg) pulmonic homograft; the aortic homograft was severely calcified, had a free regurgitation (grade 4). Angio showed a pseudoaneurysm of the left ventricular outflow tract, an abnormal origin of the coronary arteries and an occlusion of the right femoral artery. Because of the hostile chest and the very high Euroscore, the heart team decided to attempt a percutaneous valvular treatment on this grown-up congenital heart. The plan was to begin with the aortic valve which was the main problem of the patient and to do pulmonic replacement in a stage procedure. TAVI was challenging in this case because of the severe regurgitation of the degenerated homograft without any precise annular plane, with heavy calcifications and a tortuous shape; in addition, the pseudo-aneurysm of the LVOT was a risk factor for para-valvular leak at this level and the abnormal origin of coronary arteries was a potential danger for occlusion in case of suboptimal valve positioning. Finally, the right femoral artery occlusion was also an issue. Despite all these limitations, an Evolut-R 26 mm was deployed by the left femoral artery without any residual leak, improving immediately the aortic regurgitation and the hemodynamics of the patient. This second generation device is a recapturable and repositionable device allowing an evaluation of the result before final release, adjusting if needed to achieve a final accurate positioning. After a short period of recovery, the pulmonic homograft will be replaced by a Melody valve in a few weeks. In conclusion, this case illustrates how structural/interventional cardiology can offer an effective treatment in grown-up congenital heart with contra-indication for repeated complex surgeries.

#0080
THE DETERMINATION OF HYPERTENSION PERSISTENCE IN CHILDREN WHOSE AORTIC COARCTATION WERE TREATED WITH CP STENT
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Objective: Recent studies reveal that hypertension and cardiovascular diseases are more prevalent in patients with Aortic coarctation (AC), even it is treated. Our aim is to address presence of hypertension and risk for cardiovascular diseases in patients with stent applied AC by using data from ambulatory blood pressure monitorization (ABPM)/arteriography, echocardiography and radiological and biochemical evaluations.

Material and method: Twenty stented pediatric patient and 20 age- and sex-matched controls were enrolled. Physical examination findings and ECG data were recorded. Echocardiography performed...
MEDIUM TERM FOLLOW-UP OF PEDIATRIC TRANSCATHETER CAROTID ARTERY ACCESS.

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Background: Access via percutaneous carotid artery (CA) methodology is increasingly common, supplanting traditional femoral arterial access. Late incidence of CA stenosis following percutaneous access is unreported. We performed a retrospective review of patients receiving CA access to assess late CA stenosis and patency.

Methods: We reviewed all CA access cases and any follow up catheterizations that included angiography from 2012 to 2016. We reviewed access site complications, follow up ultrasound imaging of the CA, and presence of late CA stenosis on follow up angiography.

Results: During the study period, there were 48 total CA catheterizations in 41 patients – carotid access was repeated in 6 patients and twice in 1 patient. Median age at CA access was 22 days (range 1-1636) with a weight of 3.1kg (range 1.6-15.7). The majority of procedures were for ductus arteriosus (PDA) stenting (26/48; 54%) or BT shunt (14/48; 29%) interventions. Median duration of CA access was 42 minutes with time to hemostasis of 9 minutes. Acute post-cath CA ultrasound demonstrated complete luminal patency except in 2 cases (4%) where a small, non-occlusive thrombus was noted and was then treated with enoxaparin. Both patients had subsequent full

Findings: Interventricular septum thickness, diastolic LV posterior wall thickness, systolic LV posterior wall thickness, LV mass and LV mass index values were found to be significantly higher in the patient group. ABPM revealed, mean systolic pressure, daytime systolic pressure, mean arterial pressure, daytime mean arterial pressure, PWV and cardiac output values were found to be significantly higher in the patient group than control group. No significant difference was detected in aortic stiffness, distensibility and strain values between groups. CIMT was found to be significantly higher in the patient group. No significant correlation was detected among CIMT, LVMI, aix@75 and PWV values. In the comparison of 2 studies, elevation rate on months 0, 1 and 6 was 33% in the previous study while this rate was decreased to 11% in our study when assessed according to LVMass Z score.

Conclusions: It was shown that hypertension incidence and risk for cardiovascular diseases were greater than those in healthy population even AC is corrected. This suggests that AC is a part of generalized vasculopathy and that use of these measurements in follow-up of patients with AC can be helpful to predict risks.
resolution on follow up imaging within 72 hours. Late CA imaging was available in 33 patients (80%) at a median of 105 days (range 7-112) after index CA procedure with 9130 patient-days total of follow up. Among these 33 patients, there were 51 total cases of late CA follow-up imaging – all demonstrating a widely patent CA. There was no CA stenosis nor lumen irregularity noted in 33 cases (65%) with 18 cases (35%, 14 patients) demonstrating a median of 4.7% (range 2.3-12.5%) stenosis. No patient underwent intervention on the CA for vessel stenosis. No risk factors for mild CA stenosis were identified when evaluating vessel stenosis and sheath size at index procedure, interval from index procedure, weight at index procedure, or duration of CA access during index procedure.

Conclusion: Use of the CA for interventional catheterization is associated with excellent late patency with no appreciable CA stenosis noted late after the procedure. In patients with CA stenosis after catheterization, the stenosis is mild. This data supports the utilization of percutaneous carotid access during pediatric catheterization.

**#0083**

**NOVEL APPROACH TO THE DIAGNOSIS AND TREATMENT OF HEMOPTYSIS IN CHILDREN**

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**Rationale:** Hemothystis in children is an uncommon presenting symptom but can be life-threatening if massive. Cardiac catheterization and coil embolization of aorto-pulmonary collateral vessels (APCs) is uncommon in patients with hemothystis and without congenital heart disease. We present a series of 16 children with hemothystis, all of whom underwent cardiac catheterization to look for and intervene upon aorto-pulmonary vessels if found.

**Objective:** To determine the presence and size of APCs in children with hemothystis and assess resolution of symptoms after coil embolization of these vessels.

**Methods:** A detailed chart abstraction and review of bronchoscopy and cardiac catheterization images and reports was performed in this retrospective cohort study of children presenting with hemothystis in a tertiary center from January 1995 to January 2015.

**Main Results:** A total of 16 patients were identified who presented with hemothystis and underwent cardiac catheterization after bronchoscopy at our institution. The mean age was 12.6 years. Structural cardiovascular anomalies were present in 4 (25%). 13 (81%) had evidence of bleeding on bronchoscopy. Cardiac catheterization showed significant APCs (>2mm) in 9 (56%) all of which were coil embolized. 8 (89%) had complete resolution with no recurrences. 7 patients had no significant APCs. Of the 4 patients with structural cardiac anomalies, 2 had significant APCs that required coil embolization.

**Conclusions:** Aorto-pulmonary collateral vessels should be high on the differential diagnosis for pulmonary hemorrhage in an infant and besides early consultation with pediatric pulmonology, a consultation with pediatric cardiology is also warranted. Following a bronchoscopy, other imaging modalities may be helpful but cardiac catheterization would have the advantage of being able to perform interventions, specifically embolization of APCs.

**Key words:** aorto-pulmonary • collateral vessels • coil embolization • cardiac catheterization

**#0084**

**EFFICIENCY OF TRANSCATHETER PATENT FORAMEN OVALE (PFO) CLOSURE WITH HYPERION PFO OCCLUDER**

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**Introduction:** Patent Foramen Ovale (PFO) may result in paradoxical embolism event (EE). Transcatheter closure of PFO still remains an alternative and interesting treatment for selected patients. Experience with application of Hyperion PFO Occluders (similar to Amplatzer PFO Occluders) in such circumstances has been not evaluated yet.

**Materials and Methods:** 67 patients (pts) with cryptogenic EE in whom in years 2014-2016 PFO was closed percutaneously with Hyperion PFO occluder in our institution were retrospectively analyzed. There were 36 woman with mean age 41 (21-62) and 31 man with mean age 41 (19-68) years. The indications for the procedure were established by neurologist and included Transient Ischemic Attack (TIA) in 16 pts (among them 2 pts were divers), ischemic stroke in 49 pts and migraine with aura in 3 pts. Transcranial Doppler (TCD) was performed in all pts (with MES ≥2nd degree in all pts). Mean PFO tunnel length was 8 (4-16) mm. Follow up was performed 3, 6 and 12 months after the procedure and thereafter yearly. Control TCD examination was performed 6 months after the procedure and in case of residual shunt presence – 3 months afterwards again.

**Results:** The procedure was successfully completed in all patients and no procedure-related complications were observed during hospitalization. Following devices were used (accordingly to the anatomy of PFO) – PFO Hyperion Occluder 18 mm in 6 pts, 24 mm in 49 pts, 30 mm in 8 pts and 34 mm in 4 pts. Fluoroscopy time was 2,4 (0,9-22) min. During follow-up no new neurological events were observed in any pt. Control TCD results were available in 54 pts (4 pts were lost from follow-up): 48 pts had complete PFO closure and 6 pts had a residual shunt. In 1 pt paroxysmal atrial fibrillation was observed.

**Conclusions:** Transcatheter PFO closure with Hyperion devices is safe and effective procedure, however, long-term follow-up and randomized study with other devices are necessary.

**Device produced as Hyperion PFOO, Comed BV, Netherlands / SHSMA, Shanghai, China
COST OF A STANDARDIZED APPROACH TO MANAGEMENT OF PULSE LOSS FOLLOWING PEDIATRIC CARDIAC CATHETERIZATION

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Background: Pulse loss is a known complication of pediatric cardiac catheterization. A standardized approach to the diagnosis and management of post-catheterization arterial thrombosis has been shown to result in thrombus resolution in the majority of patients. The cost burden of this practice has not been reported previously and may be a perceived barrier to utilization of a clinical practice guideline (CPG) for post-catheterization pulse loss.

Methods: Single center data collected prospectively for quality improvement efforts was reviewed retrospectively. Beginning in 5/2013, all patients with post-catheterization pulse loss, identified by clinical pulse exam 1-2 hours after sheath removal, were managed according to an institutional CPG. The CPG included initial evaluation by lower extremity Doppler ultrasound, prompt initiation of anti-coagulation using low-molecular weight heparin (LMWH), and serial re-evaluation by clinical exam, Doppler ultrasound and anti-Xa levels until thrombus resolution or until 12 weeks of therapy had been completed. Charges for protocol related items: CPG follow-up visits, diagnostic imaging, medication and laboratory testing were collected.

Results: From 5/2013-2/2016, post-catheterization pulse loss occurred in 64 cases (61 patients) with a median age of 105 days (IQR 67, 154) and weight of 5.2 kg (4.3, 6.2). Incidence of pulse loss in cases with arterial access was 4.5%. By 12 weeks, restoration of pulse occurred in 92.3% of patients. Median charge per patient for pulse loss occurred in 92.3% of patients. Median charge per patient for CPG-related care was $2,672 ($1,587; $4,087) with the largest contribution coming from lower extremity ultrasound. Patients received a mean of 3±2 ultrasounds for a median radiology charge of $2,344. Pharmacy charges were minimal, with median duration of LMWH therapy of 7 days (1.23) and median charge of $82 ($25, $371). For patients with restoration of pulse prior to hospital discharge (£1 day of LMWH), median charge was $1,583 ($801, $1837). One patient on protocol was noted incidentally to have a pseudoaneurysm at 1-week follow-up, for which therapeutic thrombin injection was performed by interventional radiology. There were no bleeding events or other complications. Social burdens of chronic LMWH administration, clinical and laboratory visits for monitoring, and the financial implications of lost work/school time were not evaluated.

Conclusions: Implementation of a CPG for management of post-catheterization pulse loss results in a high-rate of pulse restoration (92%) with only modest associated charges.

COMBINED IMPACT OF A RADIATION REDUCTION INITIATIVE AND NEXT GENERATION ANGIOGRAPHIC IMAGING SYSTEM ON RADIATION DOSE IN A CONGENITAL CARDIAC CATHETERIZATION LABORATORY

Wendy Whiteside, Konstatin Averin, Russel Hirsch, Karen Minsterman, Ross Schierling, Barbara Bear, Michael Harris, Bryan H. Goldstein

Cincinnati Children’s Hospital Medical Center, Cincinnati, OH, USA

Background: ALARA (As Low As Reasonably Achievable) principles dictate best practice using the lowest possible radiation dose. Radiation reduction has been a particular focus of the congenital cardiac catheterization laboratory due to increased radiation susceptibility in children and frequent need for repeat procedures. We sought to assess the incremental impact of a radiation reduction initiative and subsequent acquisition of a next generation angiographic imaging system on radiation dose.

Methods: Beginning in 2014, a radiation reduction initiative was developed to emphasize radiation reduction within our program. This effort included equipment settings and practice changes to promote radiation lowering techniques. Subsequently, a capital upgrade to a next generation angiographic imaging system (Philips AlluraClarity) was installed in 1/2016. Single center radiation data collected prospectively for quality improvement purposes was reviewed retrospectively. The impact of these dose changes was assessed via longitudinal tracking of mean Air Kerma across the same 4 catheterization providers.

Results: From 1/2014 through 8/2016, 2083 catheterization procedures were performed at our institution. At baseline, mean Air Kerma per case was 229±476 mGy. Following initial modifications to default frame rates and ALARA-guided practice changes, we saw a 25% reduction in mean Air Kerma per case to 173±348 mGy. The installation of a next generation imaging system combined with further changes to default imaging settings generated an additional 57% reduction in mean dose per case. After 7 months with next generation equipment, mean Air Kerma per case was 49±80 mGy, a 79% reduction in dose from baseline. From 1/2014-12/2015, there were on average 5 high exposure cases (>2000 mGy) per 6-month period. Over the 7 months since equipment upgrade, there has been only one high exposure case despite similar average fluoroscopy time per case and an unchanged case-mix.

Conclusions: Changes in catheterization lab culture, practice habits, and equipment settings are effective in reducing radiation dose, but the availability of next generation imaging equipment allows for exponential dose decrease. These combined approaches result in a nearly 80% reduction in radiation dose for congenital cardiac patients.

FEASIBILITY AND VALIDITY OF PRINTING 3D HEART MODELS FROM ROTATIONAL ANGIOGRAPHY

Saar Danon, Manoj Parimi, John Buelter, Vignan Thanugundala, Sri Condoor, Wilson King

Saint Louis University, Saint Louis, MO, USA

Background: Rotational angiography (RA) has proven to be an excellent adjunctive method for evaluating congenital disease (CHD) in the cardiac cath lab, permitting acquisition of 3D datasets with excellent spatial resolution. This technique has not been routinely implemented for 3D printing. We describe our case series of models printed from RA and validate our technique.
A 5 week old male presented in congestive heart failure at 2 weeks of age with a loud continuous murmur. Echocardiogram and CT angiogram showed a fistula between the left subclavian artery and a severely dilated innominate vein. All cardiac chambers were dilated and there was systemic-level right ventricular pressure.

Procedure: Cardiac catheterization confirmed systemic pulmonary artery pressure with a Qp/Qs (effective) of 4:1. Angiography in the left subclavian artery demonstrated the fistula measuring 2.9mm at the narrowest point. The fistula was crossed from the venous side using a 4Fr Berenstein catheter and 0.018 HiTorque wire. A 6/4 Amplatzer Duct Occluder was then delivered from the venous approach placing the “hat” in the left subclavian artery and the “stem” in the innominate vein. There was no residual flow after device placement and no obstruction of the innominate vein or subclavian artery. Immediately following closure, the pulmonary artery pressure fell to 1/3 systemic. At 6 month follow-up, the infant is thriving on no medication. Echocardiogram shows no residual flow across the fistula, normal biventricular size and function and equal blood pressures in both arms.

Discussion: Congenital arteriovenous fistula (AVF) is a rare cause of congestive heart failure in infancy and mainly confined to the head and liver - only a handful of case reports describe intrathoracic AV fistulas. Once identified, transcatheter occlusion was an efficient and effective therapy leading to immediate resolution of heart failure and pulmonary hypertension.

Methods: All patients with models printed from RA were selected. RA acquisitions from a Toshiba Infinix-I system were postprocessed and printed with a Stratasys Eden 260. Two independent observers measured 5-10 points of interest on both the RA and the 3D model. Bland Altman plot was used to compare the measurements on rotational angiography to the printed model.

Results: Models were printed from RA in 5 patients (age 2 mo - 1 yr). Diagnoses included a) coronary artery aneurysm, b) Glenn shunt, c) coartcation of the aorta, d) Tetralogy of Fallot with MAPCAs, and e) pulmonary artery stenosis. There was no significant measurement difference between RA and the printed model (r= 0.990, p<0.01, Bland Altman p=0.987). There was also no significant interobserver variability. The MAPCAs model was referenced by the surgeon intraoperatively and was accurate.

Conclusions: Rotational angiography can generate highly accurate 3D models in congenital heart disease, including in small vascular structures. These models can be extremely useful in patient evaluation and management.

#0088
TRANSOCATHETER DEVICE CLOSURE OF CONGENITAL LEFT SUBCLAVIAN ARTERY TO INNOMINATE VEIN FISTULA
Jessica Nikitczuk, Barry Love
Mount Sinai Medical Center, New York, NY, USA

![3D virtual model](image1)

![3D Printed Model](image2)
**Meeting Abstracts 278**

**Daisuke Kobayashi1, Thomas Forbes1, David Nykanen2**
1Children’s Hospital of Michigan, Detroit, MI, USA
2Arnold Palmer Hospital For Children, Orlando, FL, USA

**Background:** Comparison of significant adverse event (SAE) between institutions is not straightforward because the risk associated with catheterization varies based on procedural complexity and heterogeneous patient characteristics. CCISC investigators recently proposed CRISP (Catheterization RISK Score for Pediatrics) to predict the risk of adverse events for children undergoing catheterization.

**Objective:** The study aim is to utilize CRISP Category for description of practice variation and comparison of SAE rate between institutions accounting for case volume.

**Method:** This was a retrospective study of CCISC Risk Registry. SAE rate was assessed by the funnel plots accounting for case volume in each CRISP Category. Funnel plots were depicted with 95% and 97% upper control limit of the overall mean SAE rate accounting for case volume.

**Results:** A total of 3711 pediatric patients were enrolled from 15 participating sites in 2015. SAE rates were 1.8% (13/709, Category 1), 2.0% (30/1464, Category 2), 5.3% (58/1086, Category 3), 12.0% (45/376, Category 4) and 24.2% (8/33, Category 5) [p<0.001]. The funnel plots were created for each CRISP category (Figure).

**Conclusion:** Funnel plots of SAE rate in CRISP categories were useful to show the institutional performance in a clearly visible fashion.

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**TRANSCATHETER THERAPY OF RAGHIB SYNDROME (RIGHT SUPERIOR VENA CAVA DRAINAGE INTO THE LEFT ATRIUM) UTILIZING A 3D PRINTED MODEL**

Daisuke Kobayashi, Daniel Turner, Thomas Forbes
Children’s Hospital of Michigan, Detroit, MI, USA

**Introduction:** Raghib syndrome is a rare congenital anomaly characterized by right superior vena cava (SVC) drainage into the left atrium and causes cyanosis and infection. Surgical correction is the choice of therapy and transcatheter treatment has not been described. We report a case of successful transcatheter treatment of Raghib syndrome utilizing a 3D printed model.

**Case:** A 32 year old male (104 kg) with history of brain abscess was referred for anomalous drainage of right SVC into the left atrium. There was left SVC draining into the right atrium through the coronary sinus, without a bridging vein. His baseline saturation was 90%. Cardiac magnetic resonance imaging delineated that right upper pulmonary vein drained into the right SVC. We aimed to occlude the proximal right SVC with device, achieving normal drainage of right upper pulmonary vein into the left atrium and exclusion of a right-to-left shunting, diversion of systemic venous flow from right SVC into inferior vena cava system through the azygos vein. The pre-procedural planning was augmented by the 3D Printed model. He underwent successful occlusion of proximal Right SVC with a 16 mm Amplatzer muscular VSD occluder. There was no significant increase of right SVC pressures post-device. At 6 month follow up, his saturation improved to 97%

**Conclusion:** Transcatheter therapy of Raghib syndrome was successful. Pre-procedural planning was critical, augmented by a 3D printed model in this case.

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**TRANSCATHETER CLOSURE OF VENTRICULAR SEPTAL DEFECTS: ECHOCARDIOGRAPHY BEFORE, DURING AND AFTER PROCEDURE**

M. Fernanda Biancolini, Julio C. Biancolini, Ana De Dios, Jesus Damsky Barbosa
Elizalde Children’s Hospital, Buenos Aires, Argentina

**Introduction:** The purpose of this study is to analyze the utility of echocardiography in transcatheter VSD closure.
Materials and Methods: descriptive and retrospective analysis based on the review of clinical records of patients (p) with transcatheter VSD closure during the period between May 2010 and March 2016 at Pedro de Elizalde Children’s Hospital. 51 procedures were performed in 47p. Median age was 7 years old (range 0.33 to 15y). Median weight was 26 Kg (range 4.3-83Kg); 5 procedures were done in p less than 10Kg. Sex distribution was: 28 females and 19 males.

Results: 43 procedures were successful (84.3%). 44 VSDs closed in 43 procedures. Regarding the type of VSD: 3p had residual POP VSD, the rest were native. 2p had multiple VSDs: 1p had 2 VSDs closed in the same procedure, the other needed 3 different procedures. Concerning VSD location within the septum: 20 were PM VSDs with aneurysm (46.5%); Mascul: 11 infundibular (25.5%), 6 midmucular (13.9%), 3 apical (6.9%) and 3 POP (6.9%). Respecting the size of the VSD: Median diameter of left orifice was 9.3mm, right orifice 4.7mm, and median length of 7.12mm. 12 VSDs had more than one right orifice. In regard to the relation with adjacent structures: PM VSDs had aneurysm from the septal leaflet of the tricuspid valve, 1p had mild prolapse of the right coronary cusp with partial occlusion of the VSD. 7p presented with prolapse of noncoronary cusp and 2 with right coronary cusp (1 musc infundibular, 1 POP of doubly committed VSD and the rest PM VSD). 3p revealed left ventricular-right auricular shunt, but only one more than mild (moderate). With reference to complications: 3p evidenced major complications (6.9%): 1p tricuspid stenosis (PFM); 1p hematuria with decrease in blood count (PFM); 1p, <10kg with musc infundibular VSD and no aortic prolapse or insufficiency pre procedure, had severe aortic insufficiency immediately post procedure (Amplatzer) and died. Minor complications: 11p arrhythmia, 3p needed corticosteroid treatment (2 Lifetech 1 Amplatzer); 1p, <10kg with musc infundibular VSD and no aortic prolapse or insufficiency pre procedure, had severe aortic insufficiency immediately post procedure (Amplatzer) and died. Therefore, a pre-treatment antiar - rhythmic protocol was initiated prior to the interventional procedure. Result: All lambs were followed for 6 months post implant. There was no device-associated morbidity or mortality. No significant PR and only 1 small paravalve leak (PVL) was noted from the MPA. 4/6 had a variable PVL from the RVOT that was felt to be secondary to an inadequate device size, which was confirmed at necropsy. There was tricuspid regurgitation noted in 2/6 that was associated with the delivery catheter from the RUJ. Necropsy: Smooth neotissue was overlaying the stent & valve leaflets. There was no evidence of endocarditis or thrombosis on TVP leaflets. A small area of endomyocardial fibrosis was noted in the RVOT and was attributed to the delivery catheter. Two animals had a torn tricuspid valve leaflet.

#0092
CONGENITAL CORONARY ARTERY FISTULA: PRESENTATION IN FETUS AND TRANS-CATHETER CLOSURE AFTER BIRTH
Varun Aggarwal, Venkatachalam Mulukutla, Athar Qureshi, Henri Justino
Texas Children’s Hospital, Houston, TX, USA

Congenital coronary artery fistula is a rare coronary anomaly. Most commonly, such fistulae drain into the right side of the heart or the pulmonary artery. Most children with these fistulae are asymptomatic and patients with coronary fistulae are usually diagnosed in adulthood (with symptoms like fatigue, dyspnea, angina and congestive heart failure). We describe two patients who had congenital coronary artery fistula to ventricle which was diagnosed in-utero and confirmed after delivery. One of them had congestive heart failure and the other developed ST-T segment abnormalities in the infero-lateral leads concerning for coronary steal and potential myocardial ischemia. We hereby describe transcatheter management of the large coronary to ventricle fistula. To the best of our knowledge, this is also the first description of left anterior descending and circumflex coronary ostial atresia in the setting of a large left main coronary artery to left ventricle fistula presenting in a newborn. We aim to describe the anatomy, trans-catheter closure and potential therapeutic options with concomitant limitations.

#0093
EARLY RESULTS ON THE BEIJING MED-ZENITH PT VALVE FOR PULMONARY REGURGITATION IN THE OVINE MODEL
Sharon Cheatham1, Elizabeth Clark, Chris Breuer, Joanne Chisolm1, Qingliang Zhou1, Jian Meng1, Kan Hor1, Andrew Yates1, John Cheatham1
1Nationwide Children's Hospital, Columbus, OH, USA
2Beijing Med-Zenith Medical Scientific Co., Beijing, China

Approximately 22% of all congenital heart disease (CHD) patients have anomalies of the right ventricular outflow tract (RVOT). 77% of these patients have severe pulmonary regurgitation (PR) after surgical repair. Restoring pulmonary valve function is imperative to maintaining RV function. In the past, surgical pulmonary valve replacement was the only option in these patients. The success of the Melody TVP in patients with a dysfunctional RV-PA conduit is well known. We describe the early animal results of the Med Zenith PT Valve (MZPTV), designed for the native RVOT with severe PR.

Methods: The MZPTV is composed of a symmetric nitinol frame with a porcine pericardial covering and a porcine pericardial valve inside. There are 3 valve sizes and 5 frame dimensions that are delivered through a 21 French transcatheter system. The MZPTV was implanted from the right internal jugular vein (RIJV) in 6 lambs from Nov 2015-Feb 2016: 20mm valve (n=4), 23mm (n=1), 26mm (n=1). Early experience resulted in significant ventricular arrhythmias during device delivery that precluded safe implantation. Therefore, a pre-treatment antiarrhythmic protocol was initiated prior to the interventional procedure. Results: All lambs were followed for 6 months post implant. There was no device-associated morbidity or mortality. No significant PR and only 1 small paravalve leak (PVL) was noted from the MPA. 4/6 had a variable PVL from the RVOT that was felt to be secondary to an inadequate device size, which was confirmed at necropsy. There was tricuspid regurgitation noted in 2/6 that was associated with the delivery catheter from the RUJ.
Conclusions: The MZPTV can be delivered in a safe and effective manner in the ovine model. Care must be taken to pre-treat ventricular arrhythmias that may be unique to the animal model and delivery site. The MZPTV remains competent and without significant thrombosis or infection after 6 months. PVL from the MPA is rare, while a more appropriate chosen MZPTV frame may solve PVL from the RVOT. Modification of the loading and delivery systems are underway and a clinical trial for the MZPTV is planned in China for 2017.

#0094 OUTCOMES OF PERCUTANEOUS PULMONARY VALVE IMPLANTATION – CLEVELAND CLINIC EXPERIENCE Patcharapog Suntharos1, Julia Simkowski2, Lourdes Prieto1
1 Cleveland Clinic Children’s, Cleveland, OH, USA
2 Case Western Reserve University, Cleveland, OH, USA

Background: Percutaneous pulmonary valve implantation (PPVI) has become a standard procedure performed in selected patients who have right ventricular to pulmonary artery conduit dysfunction. This procedure has been expanded to include patient with native pulmonary valve regurgitation as a result of previous surgical procedure. The aim of this study is to review our institutional experience and outcome since starting the PPVI program in 2010.

Method: Cleveland Clinic Children’s Cardiac Catheterization Database was retrospectively reviewed. All patients who underwent PPVI from 6/1/2010 to 12/31/2015 were included. Patients’ demographic and cardiac diagnosis were recorded. Clinical symptoms before and after PPVI were compared. Cardiac catheterization procedure and complications were also reviewed.

Results: Seventy-nine patients underwent PPVI during the study period with a median age of 30 years at the time of procedure (range 8 to 79 years), and 42 (53%) were male. Median follow-up time was 2.7 years (range 4 months to 6 years). The most common diagnosis was tetralogy of Fallot (49%), followed by pulmonary stenosis (24%). Majority of the patients (92%) received the Melody® Valve. The procedure was performed in RV to PA conduit in 50 (64%) patients. Eleven patients (14%) had post-procedural complications, all were considered minor. Two (2.5%) subsequently required balloon dilation of percutaneous valve for evidence of valve obstruction. Two patients (2.5%) required second PPVI, one for obstruction and one for regurgitation. No surgical reinterventions were performed in our cohort. Three patients (3.8%) developed endocarditis, all were Melody® Valve which successfully treated with long term intravenous antibiotic. Two patients died from non-PPVI related cause. Of the 78 patients for whom change in New York Heart Association Functional Classification could be determined, 30 (38%) had a decrease in classification and 48 (62%) were unchanged.

Conclusion: PPVI provides good mid-term outcome in our cohort. Similar to other studies, there was an incidence of Melody® Valve endocarditis which responded to medical management.

#0096 TRANSCATHETER LEFT ATRIAL APPENDAGE CLOSURE USING INTRACARDIAC ECHOCARDIOGRAPHIC GUIDANCE. MULTICENTER EXPERIENCE. Daniel Aguirre1, Gabriel Maluenda1, Cristian Pincetti3, Luis Perez2, Eduardo Lecanelier4, Carlos Deck4
1 Hospital San Borja Arriaran & Clinica Alemana, Santiago, Chile
2 Hospital Regional Dr. Guillermo Grant Benavente, Concepción, Chile
3 Hospital Regional Temuco, Temuco, Chile
4 Hospital Dipreca, Santiago, Chile

Most of the cases of left atrial appendage (LAA) occlusion are performed under general anesthesia using transesophageal echocardiography (TEE) guidance. ICE-guided LAA occlusion experience is limited and has been typically performed from the right atrium, pulmonary artery or coronary sinus where complete visualization of the LAA anatomy is often suboptimal. We report the feasibility, efficacy and safety of LAA occlusion using ICE guidance from the left atrium (LA).

Methods: Twenty-two patients with non-valvular atrial fibrillation, significant risk for stroke, and long-term contraindication to anticoagulation underwent LAA closure with the Amplatzer Amulet™ (St Jude Medical, Inc, St Paul, MN) device under mild sedation and ICE guidance from the left atrium. The ICE catheter and device delivery sheaths were advanced into the LA through a single transseptal puncture in all cases. Baseline patient characteristics, procedural data, effectiveness of ICE imaging, quality of occlusion, and complications were prospectively recorded. Clinical and TEE follow-up were done one to three months after the procedure. Procedural success was defined as the implantation of the closure device at the intended delivery site with adequate (grade 3 or more by angiography and ICE).

Results: Procedural success was achieved in all patients (100%). Mean age was 75±8.6 years. Previous ischaemic stroke was recorded in 9 patients (42.9%). Median CHADS2 and HASBLED scores were 3.8 (2.6) and 3.6 (4.6) respectively. Indication for LAA closure was bleeding in 5 (23.8%), ischaemic stroke under therapeutic anticoagulation in 6 (28.6%) and high bleeding risk in 1 patient(s) (2.7%). Conscious sedation was performed using midazolam and/or fentanyl as needed. The amplatzer Amulet™ device were used in all patients. Median hospital stay was 1 day. There was no procedural stroke, pericardial effusion or device embolization. No ICE or transseptal related complications were noted. At the time of writing, during the follow-up period, there were no cardioembolic events. One month after TEE showed complete LAA exclusion with no residual peri-device flow. In all cases, ICE provided adequate procedural guidance, closure assessment and LAA and device visualization, including their relation with adjacent structures.

Conclusions: Initial experience suggests LAA occlusion with the Amplatzer Amulet device using ICE guidance from the left atrium is feasible, reproducible, and safe.

#0097 SURGERY ASSOCIATED VASCULAR INJURY MAY MIMIC RESTRICTIVE SYSTEMIC-TO-PULMONARY SHUNT PHYSIOLOGY AND IS AMENABLE TO TRANSCATHETER INTERVENTION Pieter Verhoeven, Russel Hirsch, Konstantin Averin, Wendy Whiteside, Bryan Goldstein
Cincinnati Children’s Hospital, Cincinnati, OH, USA

Background: Restrictive systemic-to-pulmonary (SP) shunt physiology, typically due to anatomic shunt stenosis and/or thrombosis, is a
recognized precursor to mortality in patients with cyanotic congenital heart disease and SP shunt dependence. Surgical clamp injury with resultant vascular stenosis of a supplying artery proximal to the SP shunt may mimic restrictive shunt physiology without evidence of SP shunt pathology.

Methods: Single-center case series from 1/2012 to 7/2016. All cases with SP shunt dependent physiology involving balloon angioplasty of a systemic artery were evaluated for inclusion. Cases without angiographic stenosis of a supplying artery were excluded.

Results: Eight cases were performed in 6 patients (50% female) with median age of 94 days (IQR 71, 144) and weight of 5.0 kg (3.75, 6.2). The predominant cardiac diagnosis was hypoplastic left heart syndrome (67%). The primary interventional indication was desaturation (75%) with a median pre-cath arterial SaO2 of 71% (66, 72). The vascular obstructions presenting as restrictive shunt physiology were proximal to the shunt origin and located at the innominate artery (67%), the right subclavian artery (17%) and the right common carotid artery (17%). In 50% of cases, a second vascular stenosis was present immediately distal to the origin of the SP shunt. In all cases, balloon arterioplasty of the proximal lesion was performed with an increase in median minimal luminal diameter from 2.4 (1.5, 2.9) to 3.5 mm (3.2, 4.0), or a median increase of 52% (30, 92) over the baseline diameter. No stents were placed. No patients had SP shunt stenosis or underwent transcatheter intervention on the SP shunt or branch pulmonary arteries. Following intervention, arterial SaO2 increased to a median of 76% (76, 80). There were no procedural complications. 2 patients (33%) underwent repeat arterioplasty due to recurrent desaturation. All patients survived to stage II surgical palliation, but there were 3 mortalities (50%) by 18 months of age.

Conclusions: Surgically associated vascular injury to the arterial supply of an SP shunt can mimic restrictive shunt physiology without pathology of the SP shunt itself. Balloon arterioplasty is safe and effective in improving SP shunt and pulmonary blood flow. This cohort may be at increased risk for mortality in the medium term.

#0098
REFRAMING THE STS STAR RATING SYSTEM: A HEART CENTER PERSPECTIVE
Raghav Murthy, Kanishka Ratnayaka, Howaida El-Said, John Moore, John Lamberti

Table 1 (#0098).

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Rady Children’s Hospital, San Diego, CA, USA

Background: STS-CHSD is the largest clinical database in the world for congenital and pediatric cardiac surgery. The STS Star Rating system is based on STAT complexity categorization and mortality (increasing number = increasing complexity). Institutional cardiac catheterization volume, complexity, and outcomes are not incorporated.

Methods: We reviewed our institutional data from the 23rd harvest of STS-CHSD and included institutional cardiac catheterization data to evaluate scoring from the perspective of a comprehensive Heart Center.

Results: Inclusion of institutional cardiac catheterization data improved STAT 1 [device closure (ASD, VSD, PDA), percutaneous pulmonary valve implantation], STAT 2 [Coarctation balloon and/or stent angioplasty] and STAT 4 [RVOT stents, PDA stents, PV ballooning, adjustable PA band] reporting in this single center report.

Conclusions: Inclusion of cardiac catheterization data with traditional institutional cardiovascular surgical data may offer a more comprehensive STS Star rating system.

#0099
FEASIBILITY, SAFETY AND EFFICACY OF BALLOON DILATION OF PULMONARY ARTERY BANDS AFTER THE HYBRID PROCEDURE FOR HYPOPLASTIC LEFT HEART SYNDROME AND VARIANTS
Natália Jatene, Simone Pedra, Marcelo Jatene, Fabiana Succi, Patrícia Oliveira, Rodrigo Costa, Marcelo Ribeiro, Carlos Pedra
Hospital do Coração, São Paulo, SP, Brazil

Background: The Hybrid approach has been applied as an initial palliative procedure for neonates with hypoplastic left heart syndrome (HLHS) and variants. Precise adjustment of the pulmonary artery bands is crucial. Excessive constriction may result in severe hypoxemia in the post-operative period. Management of such patients is still unclear. We sought to determine the feasibility, safety and efficacy of balloon dilation of the pulmonary artery bands after this Hybrid procedure.
Material and Methods: This was an observational study in a cohort of infants with HLHS or variants. Data collection was retrospective. The bands were configured from 3 mm EPTFE tubes at the initial procedure. Indication for balloon dilation included persistent hypoxemia (Sats < 70%) and increased gradients on echocardiography (> 60 mmHg). Balloon dilation was performed using 2.5 to 3.5 mm angioplasty balloons. Measures of outcomes included increase in the minimal diameter of the band, increase in sats, gradient reduction across the band on echo and appropriate timing of the Norwood Glenn operation.

Results: Between March 2012 and January 2016, out of 68 neonates who underwent the Hybrid procedure, 7 were diagnosed with tight PA bands and underwent dilation at 12 band sites. The median age and weight were 60 days and 2.9 Kg, respectively. Five had HLHS (2 had variants). Three patients needed additional interventions: atrial septostomy, new ductal stenting, stenting the vertical vein and arch stenting. Arrhythmias were observed in 3 patients and minor air emboli occurred in one patient with no deleterious effects. There was no mortality. There was an increase in the minimal diameter from 0.8 ± 0.5 to 1.7±0.5 mm (p<0.001) and in the systemic saturation from 67±9 to 84±4% (p<0.001) One patient underwent a new dilation one month afterwards. In all patients the Norwood Glenn operation could be performed at an appropriate time at a median age of 6 months with no mortality. At surgery, there was a band fracture in just a single vessel. There were no lesions to adjacent vessels in any of the patients.

Conclusions: Balloon dilation of the PA bands after the Hybrid procedure was feasible, safe and effective, resulting in clinical stabilization allowing performing the Norwood Glenn operation at an appropriate time. The mechanism of dilation is probably stretching of the eptfe, which may occasionally result in fracture. No significant trauma to the adjacent vessel was observed at surgery.

#0101
TECHNICAL FACTORS ARE ASSOCIATED WITH OUTCOMES FOLLOWING RIGHT VENTRICLE DECOMPRESSION FOR NEONATES WITH PULMONARY ATRESIA AND INTACT VENTRICULAR SEPTUM: RESULTS FROM A MULTICENTER COLLABORATIVE
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Background: Transcatheter right ventricle (RV) decompression in neonates with pulmonary atresia and intact ventricular septum (PA-IVS) is technically challenging, with risk of cardiac perforation and death. Further, despite successful RV decompression, repeat intervention (ReINT) on the pulmonary valve (PV) is common. The association between technical factors during RV decompression and the risk of complications and ReINT are not well described.

Methods: Multicenter retrospective study at the 4 member centers of the Congenital Catheterization Research Collaborative (CCRC). Between 2005-15 all neonates with PA-IVS undergoing transcatheter RV decompression were included. Technical factors evaluated included: use and characteristics of radiofrequency (RF) application, maximal balloon:PV annulus ratio (BAR), infundibular diameter, and RV systolic pressure pre- and post-valvuloplasty (BPV). Primary endpoint was cardiac perforation or death; the secondary endpoint was ReINT.

Results: 93 neonates underwent RV decompression at median 3d (IQR 2-5) of age, including 61 pts via RF perforation of the PV and 32 via wire perforation. There were 12 (13%) complications with 7 occurring in the first 2 years of the 10 year study period. Complications included 8 (8.6%) cardiac perforations, of which, there were two deaths. Cardiac perforation was associated with the use of RF (p=0.047), RF duration (3.5 vs 2.0 seconds, p=0.02) and maximum RF energy (7.5 vs 5.0 Joules, p<0.01) but not with patient weight (p=0.09), PV diameter (p=0.23) or infundibular diameter (p=0.57). ReINT was performed in 36 patients including repeat BPV (n=25) or surgery (n=11). ReINT was associated with higher RV pressure (median 60 vs 50mmHg, p=0.041) and with residual valve gradient (p=0.046) post initial BPV, but not with BAR, atmospheric pressure used during BPV or the presence of a residual balloon waist during BPV. ReINT was not associated with any RV anatomic measurements, including PV annulus.

Conclusion: Technical factors surrounding transcatheter RV decompression in PA/IVS influence the risk of procedural complications but not the future need for ReINT. Cardiac perforation is associated with use of RF energy as well as RF application characteristics. Reintervention after RV decompression for PA-IVS is common, and appears related to final gradient and RV pressure after initial BPV.

ENDOCARDITIS ASSOCIATED WITH TRANSCATHETER PULMONARY VALVE IMPLANTATION: AN INSTITUTIONAL EXPERIENCE
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Objective: Transcatheter pulmonary valve (TPV) implantation has become a viable therapy for right ventricular outflow tract (RVOT) dysfunction. While rare, infective endocarditis (IE) following TPV has been reported. In this study, we describe our center’s experience with IE after Melody TPV (Medtronic, Minneapolis, MN).

Methods: Single-center retrospective review of pts who underwent TPV implantation with Melody valve from 2007-2014. IE was defined using the modified Duke’s criteria.

Results: A total of 85 pts underwent TPV implantation during the study period, of which 12 (14%) developed IE. Median time to the development of IE was 32 months (range 2-100). 2 pts had a history of IE and 2 of IV drug abuse. One pt was diagnosed with severe gingivitis prior to IE. Seven pts were on ASA at the time of IE. Compliance with SBE prophylaxis was documented in only 2 pts at the time of IE. Baseline RVOT anatomy in IE pts included homografts in 8 pts, prosthetic PV in 2 and direct RV-PA anastomosis in 2. Stenting prior to TPV implantation was performed in 8 (67%) pts. There was no significant difference in demographics, type of RVOT reconstruction, pre-stenting, and residual RVOT gradient after TPV implantation between IE and non-IE pts. Eleven pts were febrile at presentation. Eight pts (67%) presented with RVOT obstruction, 5 of which had a
vegetation in the Melody TPV on echo. Three of these 5 pts had poor outcomes: Two presented in septic shock, one was immunocompromised and died while the other underwent surgical valve replacement due to severe residual RVOT obstruction. A second death occurred in an afibrile pt with terminal heart failure who presented with multi-organ failure and IE. The other 9 pts were treated medically and none had residual RVOT obstruction after completing antibiotics. One pt developed Strep Mitis IE 2 years after TPV implantation and 5 months later had a second IE episode due to MSSA. TEE was negative both times. Most common organisms isolated were MSSA in 4 (33%) pts and Strep Mitis in 3 (25%).

Conclusion: At our institution, IE occurred in 14% of pts after Melody TPV implantation. No specific risk factors were identified. TPV dysfunction was present in 67% of cases with mortality occurring in 2 high-risk pts. One pt had their TPV explanted while the rest were successfully treated medically. Multicenter studies are needed to better understand and identify predictors of IE in this population.

#0102
PERCUTANEOUS CLOSURE OF PARAVALVULAR LEAK AND VALVE-IN-VALVE IMPLANTATION IN AORTIC REGURGITATION AFTER TRANSCATHETER AORTIC VALVE REPLACEMENT
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Paravalvular leak (PVL) is a common complication in transcatheter aortic valve replacement (TAVR) especially when significant calcification of the native valve is present resulting in underexpansion of the transcatheter valve. We present a case of a 91-year-old female patient with severe aortic stenosis who underwent right transfemoral TAVR with a 20-mm Edwards Sapien 3 Valve. After the valve was deployed, transaortic gradient normalized; however, aortography revealed severe aortic insufficiency. Elevated left ventricular end-diastolic pressure (LVEDP) and equalization of LVEDP and diastolic aortic pressure were noted on pressure tracings. Transesophageal echocardiography (TEE) demonstrated severe central and paravalvular regurgitation in the region of the non-coronary cusp. Balloon post-dilation of the transcatheter valve did not result in improvement of aortic insufficiency. The decision was then made to close the leak. Two Amplatzer Vascular Plug-II devices were used, which successfully closed the PVL. The degree of paravalvular regurgitation substantially decreased on subsequent TEE. Given central aortic regurgitation remained present, a second 20-mm Edwards Sapien 3 Valve was implanted in a valve-in-valve fashion. Final aortography and TEE showed resolution of aortic insufficiency. PVL is a known predictor of worse outcome post-TAVR and percutaneous closure of PVL should be considered as it has been shown to improve symptoms, minimize risks of hemolysis requiring blood transfusion, and is associated with improved long-term mortality.

#0103
THE SHORT-TERM RESULT OF INTRAOPERATIVE BALLOON PULMONARY VALVULOPLASTY IN VALVE-SPARING REPAIR OF TETRALOGY OF FALLOT
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Introduction: Progressive pulmonary valve regurgitation is a significant morbidity after surgical correction of tetralogy of Fallot. This study reviewed a single institution’s experience with valve-sparing repair and investigated the role of intraoperative balloon valvuloplasty.

Methods: A retrospective chart review identified 7 patients who underwent valve sparing repair of tetralogy of Fallot by intraoperative balloon pulmonary valvuloplasty. We evaluated the pulmonary valve function by echocardiography in immediate post-operative and short-term later.

Results: Among 7 patients, the number of male were 3 and median age was 136 days old. The median annulus diameter of pulmonary valve was 6.8mm (Z score -2.4) and Z score of pulmonary valve annulus was from -2.0 to -3.7. The high pressure balloon of various size were used from 8 to 12 mm and median ratio of annulus to balloon was 1.29. In immediate post-operative period, most patients had below moderate pulmonary regurgitation and stenosis. Median follow up duration from TOF repair was 22 months and most patients had mild or moderate pulmonary stenosis and regurgitation. We had 1 mortality case and she expired by endocrinology problem as adrenal dysfunction after operation.

Conclusions: In our results, the intraoperative balloon pulmonary valvuloplasty in valve-sparing repair of TOF showed feasible outcomes and long term follow up was required for more explainable results.

#0104
USING A CORE TEAM TO IMPROVE PATIENT CARE DELIVERY FOR ASD/PFO CLOSURES IN AN ADULT CARDIAC CATHETERIZATION LABORATORY.
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Background: A knowledge deficit exists at this institution in the cardiac catheterization laboratory in regards to the treatment of Adult Congenital Heart Disease (ACHD). Limited exposure to congenital cases results in a learning curve for the nursing staff and radiology technologists assisting in the procedures. Due to improvements in treatment as children, more patients are living longer, so the need for specialized adult congenital heart care has dramatically increased and will continue to do so for the foreseeable future. With the addition of Houston's only board certified Adult Congenital Heart specialist, the caseload of ACHD patients has increased at this institution.

Method: This institution's cardiac catheterization laboratory has recently implemented a core team of people that assist in these congenital heart procedures as well as briefing with the cardiologist prior to case start. Additionally, the physician meets with our clinical manager to discuss the cases coming up the following week and ensure availability of supplies that will be needed for the cases. For these purposes, we have only retrospectively assessed time to case start and total in room time in the lab for the ASD/PFO closures at this institution from 2014 to present.

Results: We reviewed the case times for ASD/PFO closures from 2014 (13), 2015 (7) and 2016 (16), specifically looking at time in
Introduction: Limb ischemia secondary to femoral arterial injury is a rare but potentially catastrophic complication of cardiac catheterization in infants. Early detection or prevention of compromised lower extremity perfusion is ideal. Near-infrared Spectroscopy (NIRS) is a well-established non-invasive method of monitoring regional oxygen saturation. The purpose of this study was to characterize intra-operative values and postoperative values of NIRS in infants undergoing cardiac catheterization involving femoral vascular access. A secondary aim was to determine if values were predictive of arterial femoral thrombus after such intervention.

Methods: Bilateral lower extremity NIRS monitoring was performed during cardiac catheterization. All infants (< 1 years old) 6kg in weight or less that underwent cardiac catheterization with femoral vascular access were eligible. Demographic and procedural data was collected. Lower extremity NIRS data was blinded as to not affect the course of the catheterizations.

Results: Data was collected for 16 consecutive infants undergoing cardiac catheterization with femoral vascular access. Cardiac diagnosis and purpose of catheterization varied between patients. Of these catheterizations, 9 were interventional. The mean lower extremity arterial-venous oxygen saturation difference (LEAVD) in the accessed leg was 21 immediately prior to catheterization, 45 immediately after sheath insertion, 52 immediately prior to sheath removal, and then 30 at the time of discharge. LEAVD at any time point was not predictive of femoral arterial thrombus.

Conclusion: Lower extremity NIRS monitoring demonstrates a widening of LEAVD during catheterization with femoral vascular access. Data on additional infants is required to determine if lower extremity NIRS monitoring in infants undergoing cardiac catheterization could be a useful and non-invasive method for early detection and potential prevention of femoral artery compromise.

Conclusion: While our results are preliminary, these findings suggest the team approach improves the quality of care of ACHD patients while undergoing cardiac catheterization and percutaneous interventions.

#0105
LOWERING REGIONAL OXYGENATION MONITORING USING NEAR INFRARED SPECTROSCOPY DURING CARDIAC CATHETERIZATION OF INFANTS.
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Methods: Bilateral lower extremity NIRS monitoring was performed during cardiac catheterization. All infants (< 1 year old) 6 kg in weight or less that underwent cardiac catheterization with femoral vascular access were eligible. Demographic and procedural data was collected. Lower extremity NIRS data was blinded as to not affect the course of the catheterizations.

Results: Data was collected for 16 consecutive infants undergoing cardiac catheterization with femoral vascular access. Cardiac diagnosis and purpose of catheterization varied between patients. Of these catheterizations, 9 were interventional. The mean lower extremity arterial-venous oxygen saturation difference (LEAVD) in the accessed leg was 21 immediately prior to catheterization, 45 immediately after sheath insertion, 52 immediately prior to sheath removal, and then 30 at the time of discharge. LEAVD at any time point was not predictive of femoral arterial thrombus.

Conclusion: Lower extremity NIRS monitoring demonstrates a widening of LEAVD during catheterization with femoral vascular access. Data on additional infants is required to determine if lower extremity NIRS monitoring in infants undergoing cardiac catheterization could be a useful and non-invasive method for early detection and potential prevention of femoral artery compromise.
Background: Children with heart disease may require repeated X-Ray cardiac catheterization procedures, are more radiosensitive, and more likely to survive to experience oncologic risks of medical radiation. Cardiac MRI is radiation-free and offers information about structure, function, and perfusion but not hemodynamics. We aim to perform radiation-free diagnostic right heart catheterization entirely using MRI guidance in an unscreened cohort of pediatric patients.

Methods and Results: We performed 45 MRI guided comprehensive transfemoral right heart catheterizations in 35 pediatric (12.7 ± 4.8 years) subjects referred for clinically indicated cardiac catheterization (post-heart transplant 37%, shunt 31%, cardiomyopathy 14%, pulmonary hypertension 12%, valvular heart disease 3%, other 3%) during an sixteen month period (3/2015 – 8/2016). Radiation-free MRI right heart catheterization attempts were all successful using passive catheters. In two subjects with septal defects, right and left heart catheterization was performed. There were no complications. One subject had six such procedures. Most subjects (51%) had undergone multiple (5.8 ± 5) previous X-Ray cardiac catheterizations. Retained thoracic surgical or transcatheter implants (40%) did not preclude successful MRI heart catheterization. During the procedure, two subjects were receiving vasopressor infusions at baseline because of poor cardiac function, and in six procedures, multiple hemodynamic conditions were tested.

Conclusions: Comprehensive MRI guided right heart catheterization was feasible and safe in this small cohort of pediatric subjects. This includes subjects with previous metallic implants, those requiring continuous vasopressor medication infusions, and those requiring pharmacologic provocation. Children requiring multiple, serial X-Ray cardiac catheterizations may benefit most from radiation sparing. This is a step toward wholly MRI guided diagnostic (right and left heart) cardiac catheterization and future MRI guided cardiac intervention.

#0108
UPDATE ON UGANDA HEART INSTITUTE 5 YEAR PROSPECTIVE PLAN TOWARD INDEPENDENT PEDIATRIC CARDIAC CATHETERIZATION
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Background: Population statistics estimate 10,000 children are born each year with congenital heart disease in Uganda. Over the past decade, Uganda Heart Institute has developed a cardiovascular surgical program. A complementary cardiac catheterization program was started in February 2012 with the aim of developing independent operation after five years of mentorship.

Methods: A biplane X-ray cardiac catheterization lab was built adjacent to a new cardiovascular operating room. Uganda Heart Institute physicians, nurses, and technologists began training with government and non-governmental organization support toward independent operation. Innovative prospective five-year independence plan includes visiting in-country clinical service and training trips, encouraging autonomy through targeted case selection (diagnostic, balloon pulmonary valvuloplasty, patent ductus arteriosus device closure), out-of-country training fellowships, and weekly telemedicine case discussion/mentorship sessions.

Results: Since inception, Uganda Heart Institute has performed 245 congenital heart disease catheterization procedures (diagnostic (85%); balloon pulmonary valvuloplasty (37%); patent ductus arteriosus device closure (102); other (21)). The number and percentage of independent Uganda Heart Institute procedures (no international provider presence) has steadily grown from 0, 0% (2012); 3, 7.8% (2013); 17, 29.8 % (2014); 55, 77.5% (2015) to 53, 82.8% (Jan-Sept 2016). In-country clinical service and training trips have been conducted by five organizations (total n = 10; World Children’s Initiative (4), Gift of Life/Chain of Hope (5)) traveling with physicians, nurses, and technologists. Two Uganda Heart Institute catheterization operators were trained at Amrita Institute of Medical Science and Research for 5 months observing and participating in 226 procedures. Weekly telemedicine conferences (n = 51 months) have facilitated dialogue and education covering clinical case selection, planning, and review. Challenges include inventory procurement, domestic hardware support, and expert support staff (anesthesia, intensive care) shortage.

Conclusions: Through mentoring, Uganda Heart Institute is progressing toward the goal of 100 annual independent cases by 2017 five years after program inception.

#0109
AMPLATZER VASCULAR PLUG II DEVICE TO OCCLUDE DIFFERENT TYPES OF PATENT DUCTUS ARTERIOSUS IN PEDIATRIC PATIENTS
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Objective: To evaluate the outcome of the Amplatzer Vascular Plug II (AVP-II) for closure of the patent ductus arteriosus (PDA) in children.

Methods: All patients undergoing transcatheter closure of PDA with AVP-II from June 2013 to November 2016 were retrospectively evaluated. In one patient device had to remove due to pulmonary hypertensive crisis. this patient excluded from study. Clinical, angiographic, and echocardiographic data were analyzed.

Results: Thirtyfive procedures were performed in 34 patients. Median age was 30 months (6–202 months) with median weight 13 kg (4.2–63). The morphological PDA classification was Type A in 14/34 (41%), Type B in 2/34 (6%), Type C in 15/34 (44%), and Type E in 3/34 (9%). The median minimum PDA diameter, maximum and length was 3.5 mm (1.5-8.3 mm), 8.2 mm (2.2-21 mm), and 13.1 mm (8.2-21 mm) respectively. The implanted device sizes were: 6 mm in 4/35 (11%), 8 mm in 14/35 (40%), 10 mm in 15/35 (43%) and 14 mm in 2/35 (6%) procedures. The implanted device was mean of
3.0 ± 1.2 times the ductus narrowest diameter and mean of 1.3 ± 0.75 times the ductus largest diameter in successful procedures. The median procedure and fluoroscopy time was 30 minutes (15-60 minutes) and 6.5 minutes (2.8-20.3 minutes). In four patients closure was performed from the arterial side. All procedures except one 100% 'in-lab' and 100% closure on post-procedural echocardiogram was achieved. No left pulmonary artery stenosis and aortic obstruction observed with a median follow-up duration of 12 months (1-40 months).

Complications: In one patient 8 mm device was embolized in to the pulmonary artery due to ductal spasm which caused underestimation of the duct and on the next day 8mm device was retrieved and a 10 mm device was implanted.

Conclusions: The AVP II seems to be an effective and safe device for PDA closure in children. It is particularly useful in type A and C ductus with adequate length and in small infants where it eliminated the risk of device related aortic obstruction.

#0110
CARDIAC CATHETERIZATION IN PAEDIATRIC PATIENTS DURING EXTRACORPOREAL MEMBRANE OXYGENATION SUPPORT
Alper Guzeltas, Ibrahim Cansaran Tanidir, Taner Kasar, Erkut Ozturk
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Background: Extracorporeal membrane oxygenation (ECMO) is a lifesaving intervention for pediatric patients with respiratory and/or cardiovascular failure. Cardiac catheterization while on ECMO was initially reserved for left atrial decompression with the balloon or blade atrial septostomy procedure, indications for ECMO support have evolved particularly by the time. Nowadays it is suggested that early catheterization after initiation of extracorporeal membrane oxygenation support in children is associated with improved survival.

Methods: Between January 2012 and October 2016, 98 patients (5.2% of the surgery patients) needed ECMO support after cardiac surgery. Diagnostic or interventional cardiac catheterization was performed in (16.3%) 16 of these 98 patients.

Results: The diagnostic catheterization was performed in 7 of the patients while invasive procedures were performed in 9 of the patients. In patients with antegrade flow, ECMO cannulas were clipped before the injections for higher quality. Median age was 6.7 months (3.3–60 months) with median weight 7.0 kg (3.7–16 kg). The median duration from ECMO support institution to catheterization was performed from the arterial side. All procedures except one 100% ‘in-lab’ and 100% closure on post-procedural echocardiogram was achieved. No left pulmonary artery stenosis and aortic obstruction observed with a median follow-up duration of 12 months (1-40 months).

Conclusions: The AVP II seems to be an effective and safe device for PDA closure in children. It is particularly useful in type A and C ductus with adequate length and in small infants where it eliminated the risk of device related aortic obstruction.

#0111
A LIFE SAVING INTERVENTION IN CRITICALLY ILL CHILDREN WITH OBSTRUCTED SUPRACARDIAC TOTAL ANOMALOUS PULMONARY VENOUS CONNECTION: VERTICAL VEIN STENTING
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Background: Obstructed total anomalous pulmonary venous connection (TAPVC) and duct dependent pulmonary flow are two cardiac pathologies which need urgent surgical intervention in pediatric cardiovascular surgery. Obstructed TAPVC has still high risk of surgical burden. Surgical mortality rate is high especially when the patients clinical condition is worse, in which preoperative stabilization by temporarily relieving the obstruction may improve outcomes in this population.

Methods: Between June 2014 and September 2016, 5 patients underwent urgent cardiac catheterization to relieve the obstruction in the vertical vein.

Results: All patients’ clinical condition was worse and intubated on the day of diagnosis. Interventional cardiac catheterization was performed within the first 24 hours after the admission. Four of the patients were newborn. Two out of five patients had isolated supracardiac obstructive TAPVC where as others had additionally right atrial isomerism, unbalanced complete atrioventricular septal defect, pulmonary stenosis in addition to and pulmonary venous hypertension in echocardiography. One of them had also duct dependent pulmonary flow.

Either 6mm or 7mm stent implantation could be achieved in all patients. In the patient with supracardiac TAPVC draining into two different parts of the SVC stenting could be performed only for one side.

Two of the patients with isolated TAPVC was operated and other one is in the waiting list.

One of the newborn patient with heterotaxy syndrome and duct dependent pulmonary flow underwent PDA stenting 5 days after the first intervention. This patient had Glenn operation and TAPVC repair when he was 6-month-old.

Remaining one patients with heterotaxy syndrome were died within two days after the procedure.

Conclusion: Stenting of the vertical vein might be an effective therapy to acutely stabilize a sick neonate with obstructed supracardiac TAPVC. Catheter intervention can be considered as part of the preoperative cardiovascular stabilization strategy for high-risk infants with obstructed supracardiac TAPVC.
Conclusion: We demonstrated a significant radiation dose reduction by implementing 3.75 fps pulse fluoroscopy rate which is the lowest possible rate. We claim that novel radiation dose reduction protocols could be easily applied without any increase in fluoroscopy time and should be applied both for patient and health care provider safety.

Table 1 (#0112).

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#0113
DESIGN AND VALIDATION OF HEMODYNAMIC CALCULATION SOFTWARE FOR DIRECT OPERATOR USE DURING CARDIAC CATHETERIZATION: THERE’S AN APP FOR THAT
Michael Ross, Jenny Zablah, Gareth Morgan, Neil Wilson
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Background: Precise and immediate hemodynamic calculations are necessary in cardiac catheterization. More complex calculations (e.g. including dissolved oxygen or using advanced models of oxygen consumption) are time consuming by hand and tend to be error-prone. Furthermore, calculations tend to be either written on-field, or performed by circulating staff that may not be familiar with the nuances of hemodynamic calculations. We sought to develop and validate a hemodynamics software application for use during cardiac catheterization.

Methods: The hemodynamics software application was developed for use with Apple™ tablets integrating accepted formulas (LaFarge...
and Miettinen) to obtain the following: cardiac output, pulmonary flow, pulmonary vascular resistance, and systemic vascular resistance all indexed to body surface area. The relationship of pulmonary flow to systemic flow and relationship of pulmonary to systemic vascular resistance was also calculated. For validation, retrospective collection of hemodynamic information was recorded and calculated in PedCath (v7.7.3) for 50 patients, age 3 months to 17 years old. Indications selected for analysis included pulmonary hypertension, atrial septal defect, and single ventricle physiology. Hemodynamic calculations were processed with the hemodynamics software and comparisons of difference between the two programs made.

Results: No significant difference between PedCath and the hemodynamics software were found. The standard error for all variables compared was <0.007.

Conclusion: We have successfully developed and validated a reliable and simple tool in the catheterization the catheterization laboratory to obtain quick and accurate hemodynamic information.

#0114
OUT WITH THE OLD AND IN WITH THE NEW: QUANTIFYING RADIATION EXPOSURE AND ITS IMPACT ON PEDIATRIC PATIENTS UNDERGOING CARDIAC CATHETERIZATION
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Objective: The application of advanced dosimetry methods to track radiation organ doses for patients undergoing cardiac catheterization.

Background: As the field of interventional cardiology in congenital heart disease continues to grow, we are becoming more aware of the potential secondary impacts of interventional treatments such as the long-term risks of radiation exposure. Currently, most institutions use metrics such as the kerma area product meter, dose at the reference point, and fluoroscopy time to track radiation. These values, while useful in charting quality control and improvements, have inherent limitations for broader application. The reporting of patient organ doses is now available to provide clinicians with an accurate method for dose tracking and the potential for patient risk modelling into the future.

Methods: A radiation dose reporting system developed at the researchers’ institution was implemented within the institution’s pediatric cardiac catheterization laboratory. Organ dose reporting was performed for a group of 40 patients within the laboratory population and a subgroup analysis was performed on 15 transplant patients undergoing cardiac catheterization in order to further evaluate clinical dose metrics.

Results: Among the entire group, the highest reported organ doses seen were in the lungs, heart, adrenals, liver and the esophagus. Among the 15 transplant patients further analysed, the average dose at the reference point for this patient group was 284 mGy. Average organ doses were 19 mGy, 12 mGy, 6 mGy, 7 mGy, and 17 mGy to the lungs, esophagus, liver, adrenals, and heart respectively. The patient with the highest doses to these 5 organs had a reported dose at the reference point of 297 mGy, which was the sixth highest in the group of 15 indicating poor correlation between these dose metrics.

Conclusions: There is poor correlation between the commonly used dose metric of dose at the reference point and the dose metric of patient organ doses. Patient organ doses provide a more accurate measure of patient radiation dose and thus may provide a more effective way of evaluating the future risk of patient radiation exposure.

#0115
INTRA-PROCEDURAL 3D ROTATIONAL ANGIOGRAPHY: SETUP TIME AND THE IMPACT OF LEARNING CURVE IN A DEDICATED “CORE TEAM” APPROACH FOR ADULT CONGENITAL HEART DISEASE INTERVENTIONS
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Background: Intra-procedural 3D Rotational angiography (3DRA) in the cardiac catheterization laboratory has been shown to add clinical value especially during complex structural and congenital heart disease interventions. Although 3D information enhances 2D fluoroscopy, some limitations that may preclude its optimal utilization include exposure to additional radiation, time needed for setup and the associated learning curve.

Methods: All adult congenital cardiac procedures performed in a single institution with intra-procedural 3DRA imaging between December 2014 and August 2016 were retrospectively reviewed. Additional 3DRA setup time was defined as the difference in the time stamps between 3DRA acquisition and preceding 2D fluoroscopic/cine acquisition. Case times and total radiation dose from fluoroscopy and 3DRA were noted for each procedure.

Results: A total of 21 patients underwent 3D rotational angiography acquisition during the study period for procedures including diagnostic pulmonary angiogram for CTEPH evaluation, trans-catheter pulmonary valve implantation, aortic coarctation and pulmonary vein stenting. The median additional setup time for 3D rotation angiography was 9.87 minutes (25th percentile: 7.87 min, 75th percentile: 19.02). Typical reasons for additional setup time include: selection of optimal imaging and contrast injection protocols, patient positioning, imaging setup and acquisition. The median radiation dose area product (DAP) was 2815.9 microGy-m2 (25th percentile: 1918, 75th percentile: 3603). There was a trend towards reduction in the set up time as operators progressed along the learning curve. Median time to case start and total in-room time were 38 minutes and 212 minutes respectively.
Residual CoA persisted in 5 animals following high pressure angioplasty (22 ATMS). The remaining 1 animal was followed as control without stent dilation and euthanized after 3 months following sham cath procedure. One planned non-survival animal was euthanized immediately after stent dilation. Two other animals died within the first 24 hours after stent dilation due to anesthesia and vascular access complications. At 3-, 6-, 9- and 12-months follow up repeat MRI, angiography and IVUS showed good stent apposition with preserved stent integrity of both BD and metal stents. Histopathology showed complete neo-endothelialization of stent material with mild-moderate inflammatory response.

Conclusions: It is feasible to deliver a novel BDS of 10-12 mm diameters to treat CoA in porcine model of CoA. The unique design of the BD stents makes it withstand the elastic forces of the aorta in resistant stenotic lesions. Further studies are needed to evaluate long term vessel/stent patency and assess risks associated with stent fragment embolization during the degradation process.

Background: 3DRA inside the catheterization laboratory is a useful intra-procedural 3D imaging tool, especially during adult congenital cardiac interventions. Development of a dedicated “core team” for adult congenital cardiac interventions, dedicated diagnostic/interventional workup and appropriate training on 3D imaging may reduce time spent on learning curve and allow more efficient use of this new imaging technology.

#0117
AMPUTATION PULMONARY ARTERY PRESSURE MONITORING USING AN IMPLANTABLE SENSOR IN PATIENTS WITH CONGENITAL HEART DISEASE

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Background: Implantable ambulatory pulmonary artery pressure (PAP) monitors are used in the management of advanced chronic heart failure (HF) in adult patients (pts) without congenital heart disease (CHD) by providing direct monitoring and remote transmission of PAP. The CardioMEMS® sensor (St. Jude Medical, Minneapolis, MN) is one such device. HF has been described after repair of complex CHD and can be responsible for clinical deterioration and repeated HF-related hospitalizations.

Figure 1 (#0116).
Methods: Retrospective descriptive review of early use of the CardioMEMS® monitoring system in pts with complex CHD. Clinical and procedural data were reviewed from 2 IRB approved studies.

Results: From 5/2015-8/2016, 8 pts (4 male) underwent catheterization with the intent to implant a CardioMEMS® device. Median age: 36.7 yr (20.8-43.8); median wt: 68.1 kg (52.4-79). Diagnoses: single ventricle s/p Fontan - 5, D-TGA s/p Mustard – 3. NYHA class: III - 7, IV – 1. Recent CHD-related admission in 7; 2 awaiting transplant. Procedural details: Standard right heart catheterization; 5 with general anesthesia, 3 with deep sedation. Vessel access for device delivery: RFV – 7, RIJ – 1; delivery sheath 11-12 Fr; device implant site: LPA – 7, RPA – 1. Concomitant procedures: LPA stent – 2, collateral vessel occlusion – 1. Successful sensor calibration and transmission of PAP in all prior to procedure completion. No procedural complications. All Fontan pts were discharged on warfarin; others were placed on clopidogrel and aspirin. In follow-up, all have successfully transmitted diagnostic quality PAP tracings. Interference with a previously implanted loop recorder was encountered in 1 pt with subsequent loop recorder explantation. There has been no evidence of thrombus formation. There have been 2 deaths in the group, unrelated to the procedure/device.

Conclusions: Implantation of the CardioMEMS® sensor is safe and feasible in adult pts with HF secondary to complex CHD. Sensor calibration is feasible and accurate in a non-pulsatile Fontan circulation and ambulatory transmitted recordings are comparable to those obtained at implant catheterization. Continued follow-up is needed to determine the impact on clinical management. This technology has the potential to provide previously unknown insight into single ventricle ambulatory physiology.

#0118
THE ROLE OF TEVAR IN THE MANAGEMENT OF ADULTS WITH COARCTATION OF THE AORTA
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Objectives: The purpose of this study was to review the utility of thoracic endovascular aortic repair (TEVAR) in adults with coarctation of the aorta.

Background: Adults with coarctation of the aorta or recoarctation may have complex anatomical substrates with recurrent obstruction or other complications including aneurysm or pseudoaneurysm formation, making traditional stenting procedures less than ideal.

Methods: Retrospective chart review was performed of all coarctation patients who had undergone TEVAR between 5/12 and 2/15 at our institution.

Results: Five patients with coarctation of the aorta underwent TEVAR. Mean age was 53 ± 13 years. Two patients had bicuspid aortic valve (one with symptomatic severe aortic stenosis) and one had previous multivessel PCI. Two patients had native coarctations and 3 were previous subclavian flap repairs of coarctation. Indications for stenting included complex native coarctation with adjacent aneurysms in 2 patients, recurrent coarctation with repair related aneurysm in 2 and pseudoaneurysm formation soon after subclavian flap repair in 1. All procedures were performed in the catheterization laboratory. Two patients underwent TEVAR alone and 3 underwent hybrid procedure combining TEVAR with vascular bypass surgery (carotid to left subclavian in 2, aorta to left common carotid in 1). Three types of endovascular stent grafts were utilized: the Gore TAG Thoracic Endoprosthesis was implanted in 2 patients, Cook Zenith TX2 in 2 and Medtronic Valiant in 1. Successful deployment was achieved in all patients. Balloon dilation of the stent graft after implantation was performed in 3 patients. Minimum aortic diameter increased from 1 ± 0.4 cm preoperatively to 1.8 ± 0.3 cm (p=<0.05). Mean blood pressure gradient between upper and lower extremities was reduced from 37 ± 18 to 7 ± 11 mmHg (p=<0.05). The number of antihypertensive medications in all patients decreased or was eliminated, mean 2.8 (range 1-6) pre to 1.4 (range 0-3) post. No major bleeding, femoral artery injury or other early procedural complications were encountered. However, 2/3 vascular bypasses had late thrombosis with transient arm ischemia in 1 and neurologic ischemia in 1.

Conclusions: Complex native or recurrent coarctation of the aorta with large aneurysm/pseudoaneurysm can be safely and effectively managed by TEVAR in selected patients. These procedures are a viable alternative to cardiac surgery.

#0119
THE USE OF BIORESORBABLE VASCULAR SCAFFOLD (ABSORB) IN RESCUE TREATMENT OF VASCULAR INJURY OF EXTERNAL ILIAC ARTERY IN TWO CHILDREN UNDERGOING INTERVENTIONAL CARDIAC CATHETERIZATION
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Bioresorbable vascular scaffold (ABSORB, Abbott Vascular, Santa Clara, California) has been recently approved as a treatment alternative for coronary artery disease. Use of bioresorbable technology has long been thought to have a theoretical advantage in children, where stented vessel may retain their growth potential. We report the use of bioresorbable stents as rescue procedure in two children with arterial vascular injury during cardiac catheterization.

Two girls, age 9 and 13 year old, who underwent interventional cardiac catheterization for a large PDA, and severe aortic stenosis, respectively, had dissection with near total occlusion of the right external iliac artery. In the first child, dissection occurred after retrieval of an embolized PDA occluder device, and the second child with withdrawal of a large balloon that failed to be withdrawn into the arterial sheath. Both children were treated with a 4 mm Absorb stent with excellent immediate results. In one child, follow up angiography done at the time of a subsequent cardiac catheterization for 2 months after the stent showed patent vessel with mild narrowing at the site of stent, no reintervention was done due to absence of clinical symptoms. Further clinical follow up 6 months after the stenting showed no symptoms with normal distal pulses. In the other child there were no symptoms of vascular occlusion, and distal pulses were normal at the time of discharge, with planned follow up.

Conclusion: We report the use of Bioresorbable stents as a rescue treatment in two children with arterial injury with good immediate results. The theoretical benefit of retaining the potential for growth in stented vessels need to be validated.
#0120
INDICATIONS, OUTCOME AND FUNDING SOURCES OF CARDIAC CATHETERIZATION IN REFUGEE CHILDREN: EXPERIENCE AT A LOW INCOME COUNTRY OF JORDAN

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Introduction: Since March 2011, hundreds of thousands of Syrian refugees crossed into Jordan. More than 650 thousand persons are registered with the United Nations Higher Commissioner for Refugees (UNHCR) in Jordan, 51% are under the age of 18 years. Demand for health care is overwhelming. Symptomatic children and young adults with suspicion of congenital heart disease were referred for evaluation at Jordan University Hospital. In this study we report our experience with cardiac catheterization in refugee patients with congenital heart disease with emphasis on types of procedures performed, outcome and funding sources.

Methods: From August 2012 to September 2016, data on refugees referred to the pediatric cardiology department at Jordan University Hospital and were found to have structural heart disease needing cardiac catheterization was recorded. Data included diagnosis, indication for catheterization, outcome, mortality and funding sources for the procedure.

Results: Of 310 refugees with heart disease evaluated in the pediatric cardiology department, cardiac catheterization was indicated in 133 patients. Funding was secured from various sources for 120 procedures in 113 patients with a median age of 4 years (5 days-30 years). Of these, 80 procedures were interventional and 40 were diagnostic. Most common intervention was closure of patent ductus arteriosus in 22 patients, pulmonary valvuloplasty in 21, coarctation angioplasty in 10, closure of ventricular septal defect in 7, closure of atrial septal defect in 6, and other miscellaneous indications in 13 patients. Three of the interventional procedures were not successful (failed aortic valvuloplasty, failed pulmonary valvuloplasty, and failed atrial septostomy for intact atrial septum). There were 10 deaths, five occurring following surgical interventions, and 5 occurred for patient waiting for surgical interventions. Mean cost of catheterization was 1050 dollars for diagnostic, and 2510 dollars for interventional procedures. Funding sources for performed procedures were individual donations in 55 (46%), nongovernment organizations in 31 (25%), UNHCR fund in 27 (23%), and patient out-of-pocket pay in 7 (6%).

Conclusion: Refugee children with heart disease who needed cardiac catheterization are more likely to be funded by charity than by United nations fund. For patients requiring surgical interventions after either a diagnostic catheterization, or palliative or failed intervention, there is a relatively high mortality, where death may occur post-operatively or while patients are waiting for funding.

#0121
CONGENITAL MULTICENTER TRIAL OF PULMONIC VALVE REGURGITATION STUDYING THE SAPIEN TRANSCATHETER HEART VALVE (COMPASSION): ONE-YEAR FOLLOW-UP.

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Background: Early safety and efficacy of the Edwards SAPIEN transcatheter heart valve (THV) in the pulmonary position has been established through a multicenter clinical trial. This study provides one-year follow-up results in an extended number of patients undergoing SAPIEN THV implantation for moderate-to-severe pulmonary regurgitation with or without stenosis.

Methods: Eligible patients were screened if body weight was greater than 35kgs and in situ conduit diameter ≥ 16mm and ≤ 24mm. Standardized implantation and follow-up protocols were used.

Results: Sixty-seven patients completed 12-month follow-up from a total of 69 implants in 81 enrolled patients. Mean weight was 71.2 ± 22.3 kgs. Indication for THV implantation was mixed (79.7%), stenosis (7.6%) regurgitation (12.7%). Twenty-two patients (31.9%) underwent implantation of 26mm valve. Patients with class I NYHA symptoms increased from 21.7% pre-procedure to 96.7% at one-year follow-up. Mean peak conduit gradient decreased from 37.7±25.4mmHg to 17.1±11.7mmHg (p<0.001). Mean estimated RV pressure decreased from 59.6±17.7mmHg to 43.4±11.9mmHg (p<0.001). Pulmonary regurgitation was mild or less in 96.7%. Freedom from device or procedure related death or reintervention was 97.1%. Two patients required early valve-in-valve reintervention, secondary to valve dysfunction. One of these developed further early valve stenosis and underwent surgical valve replacement. One patient who did not receive the SAPIEN died secondary to bacterial endocarditis involving the surgical valve. Overall freedom from endocarditis was 97.1% at one year. There were no stent fractures.

Conclusions: Transcatheter pulmonary valve replacement using the Edwards SAPIEN THV demonstrates excellent valve function and durability at one-year follow-up.

#0122
PULMONARY REPERFUSION INJURY FOLLOWING RIGHT VENTRICULAR OUTFLOW TRACT STENTING.

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Background: Pulmonary reperfusion injury (PRI) has been described in children with tetralogy of Fallot (ToF) or pulmonary atresia with multiple aortopulmonary collateral arteries undergoing unifocalization procedures to reconstruct the central pulmonary arteries. Severity has
been related to the degree of stenosis with diffuse and/or localized lung opacifications (pulmonary edema) becoming apparent on chest x-ray films within hours of having restored previously restricted pulmonary flow. We sought to determine if PRI was present following right ventricular outflow tract (RVOT) stenting. We hypothesized PRI would be related to the severity of RVOT obstruction prior stenting and would reflect on longer days of ventilation and length of hospital stay.

Methods: Prospective study including 12 consecutive children with ToF (25% female) that underwent RVOT stenting between January and July of 2016. Clinical and procedural data were reviewed. A pediatric radiologist evaluated chest x-ray films before and after intervention to determine presence and severity of PRI, as well as its impact on days of ventilation and length of hospital stay.

Results: Median age and weight were 32 months (range 1-139) and 10.8kg (1.8-22) respectively. SaO₂ increased from a mean of 71± 9.8 % to 97±6 % post RVOT stenting (p<0.0001); PaO₂ was 43±7.4mmHg and 126±94mmHg pre and post stenting respectively (p<0.001). Pre-stent RVOT diameter (mm) and PaO₂ were negatively correlated (r -0.9, p 0.037). Median time of ventilation was 5 days (range 2-8). A positive correlation was found between diameter of the pre-stent RVOT and days of ventilation (r 0.851, p 0.032) and length of hospital stay (r 0.87 p 0.024) respectively. Similarly, positive correlation was found between post-stent RVOT diameter (mm) and days of hospital stay (r 0.87 p 0.024). Regarding radiological analysis results were as follows: Out of the 12 patients, diffuse lung opacifications were classified as severe in n=5(41%); moderate n=1(8%); mild n=3(25%); none n=3(25%) within 24hs post RVOT-stenting. Pulmonary infiltrates affected the right lung in 8 (66.7%) patients. Seventy-two hours post RVOT-stent, 86(7%) of patients had complete resolution of previous abnormal findings on chest X-ray.

Conclusion: Transient PRI is present after RVOT stenting. Degree of pre-stent RVOT stenosis reflected on longer duration of ventilation and length of hospital stay.

#0123
PERCUTANEOUS PULMONARY ARTERIES REHABILITATION IN UNIVENTRICULAR PATIENTS WITH GLENN AND CRITICAL STENOSIS OR PULMONARY ARTERY BANDING
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Patients with univentricular physiology usually undergo bidirectional cavopulmonary anastomosis previous to Total Cavopulmonary Bypass. When there is high right or left pulmonary arteries pressure the Glenn univentricular physiology does not work and some times the banding of them is performed with unsuccessfully result.

Objectives: The aim of this work is to present the results of Ballon expandable Stent implantation in critical stenosis and acquired deconected pulmonary arteries in univentricular patients with Pulmonary artery banding and/or AoP Shunt.

Material and Methods: We report our experience in treatment of 4 patients with univentricular physiology ( DIU ¥LTGA with AP in 2, Ebstein Anomaly with PA and DIUV with PA with Bidirectional GLENN and LPA Banding) with severe Sytemic desaturation (69%), high Pa pressure and defuncionalized Glenn because of critical left pulmonary artery stenosis after pulmonary banding, that underwent Percutaneous Pulmonary Rehabilitation with Stent angioplasty.

There were 3 adults and one child with severe systemic desaturation even with AoP Shunt. We have performed Pulmonary angioplasty with 5 CP stent implanted in 4 patients, 2 of them covered in order to occlude the AoP Shunt, and the other nude stent. The mean diameter of stenosis was 1,8 mm for a Mean Pulmonary arteries diameter of 9 mm. After Stent implantation the mean diameter of 7,8 mm and the systemic saturation rised from 68% to 89% in all of 4 patients. The LPA MP was fram 17 mmHg to 13 mmHg.

All of patients are doing well and waiting the second stage.

Conclusion: Ballon expandable stent are useful and safe in pulmonary artery rehabilitation after BTShunt or PAB . Follow up of these patients is need to know the medium term evolution and resolution of the third stage.

#0124
COMPLETE LEFT PULMONARY ARTERY OBSTRUCTION FOLLOWING LARGE PDA DEVICE CLOSURE: SUCCESSFUL IMMEDIATE LPA STENTING TO RESTORE PULMONARY BLOOD FLOW
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Introduction: Over the last two decades, transcatheter therapy has become the first line of treatment for closure of patent ductus arteriosus (PDA). Both safety and efficacy have been established in many series and currently the procedure can be considered relatively straightforward. However, rare complications have been reported such as device embolization or device obstruction to aortic or pulmonary blood flow, particularly in small children. We present our nightmare case involving complete PDA device obstruction of the left pulmonary artery (LPA) with subsequent LPA stenting to restore left pulmonary blood flow.

Our case involves a 3 year-old, 10 kg girl with failure to thrive (< 5 th percentile). Physical exam and echocardiographic findings confirmed the presence of a large PDA and the patient was brought to the cath lab to attempt closure. Baseline hemodynamic measurements included a mean pulmonary artery pressure of 45mmHg, systolic pulmonary artery pressure was 60% systemic, Qp:Qs was >3:1 and normal PVRi (2 WU). Aortography revealed a large, type A (Krichenko) PDA, pulmonary end diameter was 9mm.

A 22/24 mm Cera™ Occluder (Lifetech Scientific) was used to close the defect. The device was deployed within the duct using standard antegrade approach. Prior release, control aortography demonstrated the device sitting within the duct without significant aortic or LPA obstruction. However, upon release, immediate proximal retraction
of the device occurred obstructing complete flow into the LPA. Initial attempts to snare and retrieve the device were unsuccessful. We then decided to restore LPA patency by stenting the vessel. In order to do so, after persistent manoeuvring we were able to pass a coronary wire into the distal portion of the LPA. Sequential coronary (1.5, 2.5 and 4mm) and conventional 4 and 7mm balloons were used to dilate and most importantly allow positioning of a long sheath. A PG1910 stent was successfully deployed achieving enough displacement of the occluder to restore LPA flow. In addition, adequate PDA occlusion was not affected by stent placement. Post-intervention, mean pulmonary artery pressure dropped to 25mmHg. There was no gradient between distal LPA and the pulmonary trunk or between ascending and descending aorta. The patient was discharged uneventfully the following morning.

**#0125**

**PATENT DUCTUS ARTERIOSUS STENTING IN NEONATES: 10 YEAR SINGLE-CENTER EXPERIENCE.**

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**Background:** Current conventional management of patients with ductal-dependent pulmonary blood flow consists of a surgically created aortopulmonary shunt (i.e. BT shunt). Experience has shown morbidity and mortality are higher when surgical shunts are performed in the neonatal period. Stenting of the patent ductus arteriosus (PDA) has become an accepted, effective and reasonably safe alternative in selected patients with ductal-dependent pulmonary circulation. We sought to report our experience with PDA stenting in neonates with ductal dependent pulmonary blood flow over the last decade.

**Methods:** Single-center retrospective review including all neonates with ductal-dependent pulmonary blood flow that underwent PDA stenting as an alternative to a surgical shunt between 2005 to 2015 at the National Institute of Cardiology Mexico City. Demographic, clinical, and interventional aspects of the procedure were reviewed. Procedure-related and early (<30 day) mortality were also included for analysis.

**Results:** PDA stenting was performed in 49 neonates (n=29 female). Median age and weight at intervention was 10 (range 1.30-3) and 3 (1.6-4kg) respectively. Clinical diagnosis included: n=22 (46%) pulmonary atresia (PA) intact ventricular septum or critical pulmonary stenosis; n=10 (20%) right atrial isomerism-PA; n=6 (12%) PA with ventricular septal defect; 6 (12%) single-ventricle physiology-PA associated with single ventricle physiology and other n=5 (10%) (i.e tetralogy of Fallot, transposition of great arteries-subPS). PDA stenting was performed through retrograde approach (via femoral artery) in n=27 (55%) cases; antegrade (venous) approach in 12 (25%) and (in more recent years) via carotid cut-down in n=10 (20%). PDA stenting was performed using different diameter bare metal coronary stents: 3mm(n=2), 3.5mm (n=29(59%)), 4mm(n=17(34%)) and 5mm(n=1). In six (12%) patients an additional stent was required to span the entire length of the ductus. O2 sats ranged preoperatively from 55± 20 to 89 ± 15 % post intervention (p< 0.0001). Six (12%) patients required an additional BT shunt due to persistence and/or recurrence of cyanosis.

Procedure-related mortality occurred in 4 (8%) patients, 2 of which occurred within the initial 5 years of establishing PDA stenting as an alternative to BT shunt in our institution. Early mortality (<30 days) was 28%.

**Conclusion:** PDA stenting is a reasonably effective alternative to surgical shunts to maintain pulmonary blood flow and increase systemic oxygenation in selected patients. In our series, procedure related mortality has decreased as we have increased our curve of experience regarding this complex technique albeit early (<30 day) mortality remains high in the neonatal period as demonstrated in our series.

**#0126**

**TRANSCATHETER DEVICE OCCLUSION OF LARGE (AND HYPERTENSIVE) PATENT DUCTUS ARTERIOSUS USING THE CERA™ PDA OCCLUDER**

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**Background:** Transcatheter device occlusion of large patent ductus arteriosus (PDA) is no simple task. Traditionally, non conventional double-disk devices (atrial and ventricular septal occluders) have been used provided they are available in large diameters albeit their shape is not specifically designed to accommodate into most of the frequently encountered PDA morphologies. In this setting, the Cera™ PDA occluder (Lifetech, Scientific) is similar to the standard Amplatzer Duct Occluder but has a broader range of diameters (up to a maximum of 24mm) making it an attractive device to close large defects.

**Methods:** Single centre, retrospective analysis of 39 consecutive patients (female N=27, 69%) with large PDA that underwent closure using the Cera™ PDA occluder between May 2012 and November 2015. Standard implantation technique was performed in all cases. Careful hemodynamic assessment was mandatory. Vasorreactivity test (100% O2 and/or iloprost) and/or temporary PDA balloon occlusion was performed if moderate to severe pulmonary arterial hypertension was present and closure considered if pulmonary vascular resistance was ≤ 6 Wood units. Continuous variables were expressed as median and range or mean and standard deviation were appropriate, while categorical data were expressed as numbers and percentages. Paired t-test was used for post-procedure hemodynamic observations.

**Results:** Median pulmonary and aortic end diameters were 10(6.8-12.7) and 17.5(10-27) respectively. Mean pulmonary systolic artery pressure (PSAP) was 69±21mmHg with a Qp:Qs ratio of 3.6 ±1.7 to 1. Successful closure was achieved in 38 (97%) cases. Mean PSAP decreased to 47±14mmHg (p<0.001), Device embolization occurred in 26(5%) cases. The first patient had the original device snared and retrieved uneventfully followed by successful deployment of a larger device. The other patient required surgical retrieval of the device and PDA closure. Follow-up was available in all patients with a median follow-up of 24(9,61) months. Currently, 30(79%) patients have completed 1-year follow-up. In this group, mean PSAP was 37±13mmHg on echo. No significant left pulmonary artery obstruction (≥10mmHg) has been encountered.

**Conclusion:** The Cera™ PDA occluder is safe and effective for closure of large and hypertensive defects.
Pulmonary vein obstruction is poorly tolerated in patients with Fontan palliation and total anomalous pulmonary venous return (TAPVR). A 19 year old male with heterotaxy and single ventricle, bilateral superior vena cavae (SVC) and TAPVR to the SVC- right atrium (RA) junction who was had previously undergone extracardiac non-fenestrated Fontan was transferred to our institution after being admitted with cyanosis and chronic pleural effusions. Diagnostic cardiac catheterization revealed multiple veno-atrial collaterals, elevated Fontan pressures at 30mmHg, and elevated pulmonary capillary wedge pressures (PCWP) of 24mmHg compared to end diastolic pressure of 9mmHg. Transeosophageal echocardiography (TEE) showed pulmonary vein stenosis localized to the entrance of the pulmonary venous confluence to the SVC. Due to the perceived difficulty of an intervention from the femoral vein approach, a hybrid approach was performed in a single plane hybrid operating room. The right atrium was exposed through a repeat median sternotomy incision and a pledgetted pursestring suture was placed in the anterior wall. A 14 French short sheath was inserted into the RA, and using TEE and fluoroscopic guidance, a wire was easily manipulated across the pulmonary vein confluence into the right lower pulmonary vein. Balloon compliance testing was performed with a 16mm x 3cm Tyshak II balloon demonstrating a discrete waist that resolved easily with inflation but returned upon deflation. A 26mm EV3 Mega LD stent was hand crimped on a 20mm Numed BIB balloon and deployed across the stenosis. There was unobstructed flow and no significant gradient by TEE or catheter pullback pressure following stent deployment. The patient recovered well from the procedure. All chest tubes were removed by postoperative day (POD) 8, and he was discharged home on POD 12 with oxygen saturations in the low 90’s in room air. This case demonstrates the utility of hybrid procedures in treating complex obstructions in high risk single ventricle patients.

#0128
ELECTIVE PALLIATION WITH SERIAL CATHETER-BASED INTERVENTIONS AS A BRIDGE TO SURGICAL REPAIR IN PREMATURE INFANTS WITH OBSTRUCTED TOTAL ANOMALOUS PULMONARY VENOUS RETURN
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Premature infants born with obstructed total anomalous pulmonary venous return (TAPVR) are often not considered candidates for surgery and therefore pose a significant treatment dilemma. We describe the course of two premature infants with obstructed TAPVR who were palliated with serial catheter based interventions. Case One: A 1.9kg infant, born at 32 weeks gestational age, was diagnosed with infra-diaphragmatic TAPVR and mild obstruction at the level of the ductus venosus. Although he could be stabilized initially with aggressive respiratory support, serial echocardiograms demonstrated increasing obstruction. On day of life (DOL) 10, stenting of his ductus venosus was performed by a right internal jugular vein approach with two 4mm Medtronic Integrity stents. He was extubated after the procedure and remained stable for several weeks. He developed progressive hypoxemia and pulmonary edema associated with increasing gradient across the stents by echocardiogram. The stents were dilated with a 5mm Bard Dorado balloon with relief of the gradient and this allowed for stabilization until he underwent surgical repair at DOL 60 and a weight of 2.8kg. He had an uneventful postoperative course and was ultimately discharged home. Case Two: A 1.4kg infant, born at 30 weeks gestation developed respiratory distress in the immediate post-natal period and was diagnosed with supra-cardiac TAPVR draining to the left innominate vein. Due to progressive obstruction, stenting of his vertical vein was performed at DOL 41 with a 4mm Medtronic Integrity stent through femoral vein approach with relief of the gradient. He developed progressive obstruction across his stent, and he was taken back to the catheterization lab where a 5mm Cook Formula stent was placed within the previous stent and post dilated with a 6mm Sterling balloon providing relief of the gradient. This allowed stabilization to surgical repair at DOL 79 and a weight of 2.9kg. He had an uneventful postoperative course and was ultimately discharged home. Serial catheter-based palliative interventions in premature infants with TAPVR should be considered as a management option that allows them to achieve an acceptable weight and maturity for surgical repair.
Implanted valve diameters were 26 (n=1), 28 (n=1), 30 (n=2) and 32 mm (n=1). Four valves were 30 mm in length and one was 25 mm. No pt had a significant RVOT gradient or PR after the procedure. Mean length of stay was 3 days (1-5). Two pts had low-grade fever after the implantation with negative blood cultures. In a mean follow-up of 3.4 months (1-5) PR is grade 0 in all pts, mean Doppler systolic peak gradient across the valve was 15.8 mm Hg (14-19), 1 pt developed a sustained ventricular tachycardia 18 days after implantation requiring cardioversion and anti-arrhythmic medication and there was 1 case of stent strut fracture on fluoroscopy with normal valve function.

Conclusions: PPVR using the Venus P-Valve® in pts with enlarged RVOT was feasible, safe and effective in this preliminary experience. Immediate pulmonary competence was seen. Further follow-up studies are needed to evaluate the long-term valve performance.

**#0131
COMPREHENSIVE ASSESSMENT OF CORONARY ARTERY COMPRESSION DURING TRANSCATHETER PULMONARY VALVE PLACEMENT: THE ROLE OF DYNAMIC PROJECTION CORONARY ARTERY ASSESSMENT**


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Background: Coronary Artery (CA) compression is a known devastating complication of Transcatheter Pulmonary Valve Placement (TPVP). Fixed Projection Imaging (FPI) is limited in visualizing the complex relative anatomy of the CAs and the Right Ventricular Outflow Tract (RVOT). We have developed a strategy involving selective CA angiography during conduit dilation for CA compression testing (CACT) using Dynamic Projection Coronary Artery Assessment (DPCAA) (3-dimensional rotational angiography, CA Swing angiography, or manual rotation CA angiography). We sought to compare these DPCAA to FPI in the setting of CACT prior to TPVR.

Methods: Retrospective study from subjects who have undergone CACT during from 2010 to 2015. Demographic, echocardiographic, hemodynamic and angiographic data were reviewed. Patients who underwent selective CA angiography during CACT were included for comparison. Comparison of the number of angiograms needed for CACT was performed between those who underwent FPI to those with DPCAA.

Results: There were 132 subjects with intent for TPVP. 102 subjects met inclusion criteria with a median age of 22 years (IQR: 14.3 – 29), weight of 60.5kg. (IQR: 51.4-72.1), with 36% female gender. The indication for TPVP was: pulmonary stenosis 30(29%), pulmonary insufficiency 22(22%), and mixed pulmonary stenosis/pulmonary insufficiency 50(49%). Nine subjects (9%) were found to have coronary artery compression precluding TPVP. Sixty-one subjects underwent selective coronary artery assessment. Of these, 16 subjects (15%) underwent DPCAA. There was a significant decrease in the number angiograms needed in patients who underwent DPCAA vs. FPI (3.60 vs. 4.74 angiograms, p-Value: <0.01).

Conclusion: Dynamic Projection Coronary Artery Assessment can aid in decreasing the number of angiograms needed to diagnose coronary artery compression during Transcatheter Pulmonary Valve Placement.

**#0132
FEASIBILITY AND INITIAL OUTCOME OF TRANSCATHETER CLOSURE OF PERIMEMBRANOUS VENTRICULAR SEPTAL DEFECT WITH DIFFERENT DEVICES**

**Manish Shrestha Shrestha**, **Worakan Promphan**

As the medium-term evolution of the uncorrected defect, patients have a very bad prognosis. Previously there was only the possibility of closure by surgery, but now the closure in the cath lab is a reality.
**Background:** Transcatheter closure of perimembranous ventricular septal defect (pmVSD) remains controversial due to its association with higher incidence of complete heart block (CHB). Various devices are being developed to minimize the procedure-related complications. The aim of this study is to analyse the feasibility of closing pmVSD with different devices by a percutaneous approach and determining the initial six-month outcome of the procedure.

**Methods:** A retrospective longitudinal cohort study was conducted in all consecutive patients who underwent transcatheter closure of pmVSD from September 2009 to March 2015 in two major cardiac centers in Thailand. The enrolled subjects were followed up for six months with electrocardiogram and transthoracic echocardiogram.

**Results:** 146 patients underwent transcatheter closure of pmVSD. The median age at intervention was 7.5 years (ranging from 9 months to 28 years). Seventy-five patients (51.4%) were male. The median weight at intervention was 22.5 kg (ranging from 6.4 to 100 kg). Thirteen different devices were used to occlude pmVSD. Transcatheter pmVSD closure was successfully performed in 142 cases (97.2%). There were two cases (1.4%) of new onset significant aortic regurgitation (AR) and one case of CHB (0.7%). During a six-month follow-up, there was no mortality although 21 patients had minor adverse events. At immediate follow-up, 32% had mild to moderate residual shunt and 20% had trivial to mild AR. At 6-month follow-up, 11% had mild residual shunt and 14% had trivial to mild AR.

**Conclusion:** With proper case selection, good expertise and judicious use of various devices with respect to anatomic details of pmVSD, transcatheter closure is feasible with promising outcomes.

**Key words:** complete heart block • device • perimembranous VSD • and transcatheter closure

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**#0133**

**PERCUTANEOUS COIL CLOSURE OF LARGE LEFT CORONARY ARTERY FISTULA IN A 48 Y.O. WITH NORMALIZATION OF SEVERELY DEPRESSED LV WALL MOTION.**

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**Introduction:** Coronary artery fistulas are a rare cardiac defect with an incidence of 0.002% and account for 0.2-0.4% of all congenital heart defects. They most commonly arise from the left anterior descending or right coronary artery with the majority of fistulas draining to the right-sided heart structures. Small case series report approximately 4% drain to the left ventricle. The majority of fistulas are small and asymptomatic. We could not find in a review of the literature whether patients presenting with severe depression of the LV wall motion could be reversible after closure. We describe a case where the EF normalized on ECHO the following day and is normal on follow up to 6 months.

**Methods:** A 48 y.o. male presented with dyspnea; exertional fatigue and exercise intolerance and was found to be in atrial fibrillation and had an EF of 25%. He was placed on a life vest; Eliquis (Bristol-Myers Sqibb) and limited activity restrictions. His atrial fibrillation was rate controlled with no improvement in his EF. He therefore had a cardiac cath, which demonstrated a moderate sized tortuous coronary artery fistula arising off of the distal LAD, which appeared to be draining to the LV. He was therefore referred for percutaneous closure.

Selective left coronary artery injections revealed a very tortuous fistula off the LAD measuring 5 x 3.9mm coursing across the RV apex with an intramural course with drainage to the LV with narrowing to 3.5mm. Balloon test occlusion was performed without ST segment changes. The fistula was then closed with a 6mm × 20 cm Interlock-18 Coil (Boston Scientific), followed by a 5mm × 8 cm Interlock-18 Coil with complete occlusion. His Eliquis was continued.

**Results:** Pre-cath ECHO EF was 25%. Post-cath ECHO EF the following day was 64%. The patient’s symptoms have resolved and patient is now able to exercise again. He had a routine screening cath performed 4 months after the procedure which demonstrated complete closure; no proximal extension of thrombus and improved filling of proximal branch vessels of the LAD.

**Conclusion:** This is only a single case report. However, our experience would suggest that poor LV wall motion secondary to a coronary artery fistula may be reversed with closure even in the adult patient with presumed long standing shunting. In addition, percutaneous coil occlusion is a possible, safe treatment option even when draining the LV. Long term anticoagulation strategies are not well established.

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**#0134**

**HYBRID APPROACH TO PREMATURE INFANTS WITH COMPLEX CONGENITAL HEART DISEASE OF JEHOVAH’S WITNESS PARENTS PURSUING BLOODLESS TREATMENT OPTIONS.**

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**Introduction:** There are very few programs that offer intervention and there are limited options for neonates born with complex congenital heart disease whose parents are Jehovah’s Witness and are pursuing bloodless options. We present two patients that underwent a percutaneous intervention in the neonatal period with subsequent bloodless full surgical repair after one year of age.

**Methods/Results:** Case #1: 4 month former 32 weeker with birth weight of 1.8kg infant with pulmonary atresia, VSD, hypoplastic confluent branch PAs and stenotic MAPCAs with O2 sats of 68%. She was referred for possible percutaneous intervention. She underwent...
perforation of the 3.5 mm pulmonary valve into a 1.8mm MPA and placement of a 4mm Formula 418 Stent (Cook Medical) in the RVOT and MPA with confluent branch PAS measuring 1.5mm each. Her O2 sats increased to 88%. She underwent a second cath at 8 months of age to redilate the RVOT stent to 5mm. She underwent a successful bloodless repair at 14 months of age with RV-PA homograph/VSD closure.

Case #2: 36 weeker with birth weight of 2.4kg with Hypoplastic Aortic Arch and VSD. Patient underwent placement of a 4mm X 8mm Rebel Stent (Boston Scientific) in the PDA and placement of branch PA stents at 2 weeks of life. He had the stent redilated at 6 months of age to 5mm. He underwent a successful bloodless arch repair and VSD closure at 17 months of age.

Conclusion: Both patients had 2 caths and one full repair on bypass without receiving any blood products. This percutaneous hybrid approach provides a safe alternative and allows patients to delay the first surgery until after a year of life and at least 9 kg to be able to undergo a bloodless surgical repair.

#0135
BALLOON-EXPANDABLE COVERED STENT IMPLANTATION FOR TREATMENT OF A TRAUMATIC PEDIATRIC PULMONARY ARTERY PSEUODANEURYSM
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Traumatic pseudo-aneurysms of the aorta are well described consequences of motor vehicle collisions in adults, but pseudo-aneurysms of the pulmonary arteries are more often attributed to infectious etiologies, such as necrotizing pneumonia, bacterial endocarditis, tuberculosis, aspergillus, or vasculitis. Traumatic pseudo-aneurysms in the pediatric population are extremely rare, and often require a novel approach with considerations for continued somatic growth of the vessel and minimally invasive techniques. Here, we detail the case of an 8-year-old male passenger in a motor vehicle collision, whose trauma led to the formation of a pulmonary artery pseudoaneurysm. Given the poor candidacy for open surgical repair, an endovascular approach was performed using the NuMed Cheatham Platinum (CP) covered balloon-expandable stent (NuMed, Inc., Hopkinton, New York). The advantage of the use of a balloon expandable CP covered stent in the pediatric setting is the stent’s ability for further dilation on subsequent evaluations if necessary and with somatic growth. The first CP covered stent was placed over the distal pseudo-aneurysm in the left pulmonary artery (LPA), and a second CP covered stent was coupled to the first stent extending into the main pulmonary artery (MPA) and “flowered” to obtain apposition to the MPA wall without dilating the LPA. Subsequent CT imaging demonstrated stable stent placement without evidence of complication or a distal filling defect.

#0136
15 YEARS OF EXPERIENCE WITH PERCUTANEOUS TRANSCATHETER CLOSURE OF ATRIAL SEPTAL DEFECTS IN PATIENTS LESS THAN 12 YEARS OLD, MEXICO CITY.

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The first transcatheter closure of an Atrial Septal Defect (ASD) in our NIH was in 2000, at the beginning all patients under 12 years were operated for the closure of the ASD. The objective of our report is to present the evolution in this group of patients, the change of the indications of transcatheter closure (TC) vs surgery, the percentage of success cases, and the risk factors for complications in this group of patients. We examined the outcome variables through analysis of the record files, comparing the patients who underwent catheterization to those who underwent open surgery for the closure of the ASD.

We present a total of 44 patients under 12 years from 2000 to 2015. The proportion of TC/Surgery for closure of ASD changed significantly during the years, and the most representative years are 2006 when we closed 13 ASD by surgery vs 2 by TC, and 2014 with a total of 12 ASD closed by TC vs 2 by surgery.

Total of 44 patients, 10 (23%) cases were classified as complex, 97% had a successful closure. In the first 24 hours they present mild flow thru the device in 10 patients (27%) and moderate in 1 patient (2.7%), at 6 months 100% had a total closure of the defect. A total of 3 cases had complications (6.8%) of which 2 were rhythm disorders (4.4%), 1 case the device embolized (2.2%). Patients <20kg were more likely to had a complex ASD.

Percutaneous closure of atrial septal defect is a safe procedure in patients under 12 years. This study demonstrates the natural evolution in the expertise to perform a procedure, as cases solved by percutaneous closure of the ASD in 2015 would have underwent surgical closure in 2000. We still need to acquire experience in our center for percutaneous closure in patients under 4 years old (16 kg).

#0137
RISK FACTORS FOR REINTERVENTION IN CHILDREN UNDERGOING TRANSCATHETER ATRIAL SEPTAL INTERVENTIONS FOR ATRIAL DECOMPRESSION AND MIXING
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Objectives: Catheter-based atrial septal interventions (ASI) to create or enlarge atrial septal defects (ASD) are performed for left atrial decompression (LAD), right atrial decompression (RAD) or to enhance atrial level mixing or streaming in infants and children. Occasionally, reintervention (RI) is necessary due to restriction of the ASD. The goal of our study was to identify risk factors for RI based on patient factors and technical approaches.

Methods: A retrospective chart review was performed on all patients that underwent ASI between 2005-15, including: balloon atrial septostomy (BAS), septal stent, or static balloon dilation (SBD). Infants with transposition of the great arteries were excluded. Data collected
#0139 PERCUTANEOUS RECONSTRUCTION OF INTERRUPTED AORTIC ARCH IN ADOLESCENTS AND ADULTS

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Percutaneous stenting seems to be an attractive & preferred alternative to surgical therapy for treatment of severe aortic coarctation in the adolescents & young adults. However such procedures are challenging. In the last 2 years, five of all patients who were referred to our center as a cases of coarctation of aorta were found to have interrupted aortic arch during cardiac catheterization. The clinical examination & CT-angiogram were suggestive of severe coarctation of the aorta. Cardiac angiography showed an interrupted aortic arch, just distal to the origin of the left subclavian artery with a gap ranging from 5 – 8 mm between the proximal & distal segments.

Patients & Methods: Right axillary & femoral arterial access were obtained. Antergrade & retrograde aortograms were done in diagonally apposite projections to analyze the relation between proximal & distal segments. A straight end hole catheter was crossed in the distal segment & another pigtail or straight end hole catheter was firmly engaged in the floor of the proximal segment to perform an angiogram to visualize both segments simultaneously. In some cases BMW 0.014 Fr. x 182 mm guide wire can be crossed from the proximal segment to the distal one, then a guide wire 0.021 Fr. or 0.025 Fr. x 150 mm crossed antegradely to enlarge the hole in the membrane. If this method was unsuccessful, the proximal gap was perforated antegrade by aprogress guide wire 0.014 x 182 mm or pilot guide wire 100-150 x 182 mm using proximal & distal aortogram as a road map thereafter there are 2 methods to dilate the tiny hole either by inflating a PTCA balloon 4 x 20 mm (sprinter legenb) up to 16 bar or by sequential dilatation of the hole with an incremental sized-dilators until 12- 14 Fr. Cook sheath crosses to the proximal segment where Z-mid or BIB-balloon mounted covered CP- stent progressed over the Amplatz guide wire 0.035 Fr x260 mm was inflated at the site of interrupted segment. After stent inflation, pressures were taken in the proximal & distal segments with post-stenting aortogram showing good antegrade flow without any complications.

Results: All those patients underwent successful procedures with no residual pressure gradient or procedure- related complications after CP- stenting of the interrupted segment. At 6 months follow up, they were asymptomatic & their blood pressure was controlled.

Conclusion: Although transcatheter reconstruction of interrupted aortic arch is technically difficult & challenging but it is feasible & associated with high success rate. Different techniques & tricks positively influence the outcome.
**#0141**
**IS A LEFT-SIDED HEMIAZYGOS CONTINUATION OF THE INFERIOR CAVAL VEIN AND MÉSCARDIA A CONTRAINDICATION FOR A PERCUTANEOUS PULMONARY VALVE IMPLANTATION? A CASE DESCRIPTION.**

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**Introduction:** Percutaneous pulmonary valve implantation (PPVI) is today an established intervention. In the majority of the cases is preferred the trans femoral approach and only in special circumstances is used the right trans jugular approach.

**Methods:** We describe PPVI by a 14 year old female with tetralogy of Fallot, mesocardia and Hemiazygos continuation of the inferior caval vein, left sided superior caval vein (LSVC) draining in the coronary sinus. At 4 months of age underwent for a ventricular septal defect closure, with homograft insertion from the right ventricle to the pulmonary artery, patch augmentation of the left pulmonary artery (LPA) and creation of an atrial communication (ASD). Thereafter she underwent numerous catheterizations due to a stenosis of the LPA and at the age of 5 she underwent device closure of the ASD.

At the age of 12 due to a Homograft stenosis she underwent for a balloon dilatation and one year later for a stent implantation (covered CP 8Z34 of NuMED Inc. over a 20mm Z MED II Balloon of NuMED Inc.) in the Homograft as a preparation for a future PPVI. Prior to the PPVI she underwent a second stent implantation into the LPA (AndraStent XL 30mm from Andramed over 15 mm Z MED II Balloon) and in the Homograft distal to the first one (AndraStent XXL 39 mm over a Z MED II Balloon 22 mm). Finally, a 22 mm Melody® Valve was implanted through the Hemiazygos vein.

A pre formed extra stiff guide wire (Lunderquist, Cook Medical inc.) was placed through the Hemiazygos -LSVC draining in the LPA and formed a "U" type root. The large delivery sheath of the Melody Ensemble kept a smooth undistorted shape, in order to deliver the valve. The valve was positioned precise in the landing zone.

The procedure was performed under deep sedation according to our standard protocol. The duration of the procedure was 172 min and the radiation time was 24.9 min.

**Discussion:** Reports in the literature support the root of jugular vein as feasible and safe for PPVI. There are no reports as far to our best knowledge of literature, which describe a PPVI through a Hemiazygos-LSVC-CS pathway.

The "U" type configuration, which is the same configuration as for the access from the right superior caval vein was crucial in order to deliver the Ensemble in the proper position. To achieve this, the extra stiff exchange wire had to be manually shaped according to the underlying anatomy to enable a smooth and safe placement of the long sheaths and delivery system. This configuration provided us the proper support and accessibility for all prior interventions as for the PPVI. The duration of the procedure and the radiation time was comparable to those reported in the literature.

**Conclusion:** Based on this unique experience, PPVI is possible even in patients with unusual anatomy. Establishing a "U" shape form of the guide wire according to the underlying anatomy is a helpful tool to enable a safe positioning of the valve.

**Background:** Patients with functional aortic interruption of the descending thoracic aorta at the isthmus due to severe coarctation are extremely rare and frequently solve by surgery.

**Case presentation:** We describe case with radial access (stiff guide) because the impossibility to reach the ascending aorta from the femoral arteries. After that we use an arterio-arterial loop with the help of a Snare catheter, and a successful percutaneous reconstruction using a covered stent (CP).

**Conclusions:** This report is an attempt to highlight the role of minimal invasive approach in the management of a severe coarctation of the aorta and the utility of use the radial access in pediatric patients to avoid morbidity and mortality associated with more invasive procedures.

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**#0143**
**VALEO STENT IMPLANTATION FOR RECURRENT AORTIC ARCH OBSTRUCTION FOLLOWING THE NORWOOD OR DAMUS-KAYE-STANSEL PROCEDURE.**

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**Background:** Stent implantation for recoarctation of the aorta following Norwood or Damus-Kaye-Stansel (DKS) procedure is well established as an effective treatment.

**Methods:** Evaluate the prevalence and outcome of stent implantation for recoarctation in children following Norwood or DKS procedure over the last decade at a single national cardiology centre. In particular we sought to assess the performance of the Valeo stent (Bard Peripheral Vascular, Tempe AZ) in this setting.

**Results:** Of 114 children who underwent Norwood procedure or DKS between January 2003 and March 2013, 80 patients survived the peri-operative period. Of these fifteen children underwent stent implantation for recoarctation. A Valeo stent was employed in 11 children, a Palmaz Genesis stent in 2 patients, a MultiLink stent in one child, and a Jo stent/covered stent in one child. The only predictive factor of need for stent placement was previous angioplasty (p value<0.01, Chi square 11.5). Eight of these patients had previous balloon angioplasty.

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The Valeo stent was inserted at a median age = 3.8 months (IQR 3.8 – 10.2 months). The median reduction in aortic arch gradient was 24 mmHg (IQR = 14.5 – 29 mmHg) [p<0.01]. The median increase in aortic diameter = 3.1 mm (IQR 2.7 – 4.2 mm) [p<0.01] and the median increase in coarctation index = 0.40 (IQR = 0.33 – 0.46) [p<0.01]. There were no procedural deaths but stent migration occurred in one child. One patient (1/11) died during follow-up due to poor systemic ventricular function. Six patients required redilation of the stent with no complications.

Conclusions: The prevalence of stent placement for recoarctation following Norwood/DKS operation was 15%. Recoarctation was successfully addressed by percutaneous insertion of a Valeo stent. Due to its low profile and facility to re-dilate to much larger diameters, the Valeo stent is a useful addition to the interventionalist’s armamentarium and in particular for relief of recoarctation of aorta in univentricular infants.

#0144
RELIEF OF OBSTRUCTED NATIVE PARTIAL ANOMALOUS PULMONARY VENOUS CONNECTIONS WITH FORMULA STENT IMPLANTATION
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Background: Transcatheter relief of total anomalous pulmonary venous connections (TAPVC) has been employed to stabilise patients pre-operatively. There are a number of encouraging case reports of stent implantation for obstructed TAPVC as an initial intervention but none relating to partial APVC (PAPVC).

Report: Ex 32 week gestation infant unable to wean off ventilatory and inotropic support. Echocardiography demonstrated PAPVC of the left upper pulmonary vein to the innominate vein, a small secundum atrial septal defect with bidirectional shunting, mild right heart dilatation, moderate right ventricular hypertrophy but well preserved biventricular systolic function (left ventricular ejection fraction = 86%). There was moderate pulmonary regurgitation and mild tricuspid regurgitation. The development of left sided pulmonary oedema occasioned a CT angiogram which suggested vertical vein stenosis with no apparent extrinsic compression.

At eight weeks (weight 3.3kg) cardiac catheterisation was performed which confirmed ¾ systemic pulmonary arterial pressures and a severe discrete stenosis in the vertical vein receiving the left upper and part of the left lower lobe pulmonary veins (Figure 1). The peak pressure gradient across the stenosis = 9 mmHg.

A pre-mounted Formula 414, 6mm×12mm (Cook Medical, Bloomington, IN) stent was placed across the stenosis, over a 0.014" Asahi Grand Slam wire (Abbott, Illinois, USA) and inflated to 22 ATM. There was excellent angiographic relief of the stenosis with a 1 mmHg pressure gradient across the stent. The procedure was well tolerated.

Conclusion: Obstructed PAPVC may potentiate pulmonary hypertension necessitating intervention. Endovascular stent implantation for obstructed supracardiac PAPVC is technically straightforward, but the stenosis is often resistant, and may provide effective relief of the stenosis with reduction in pulmonary artery pressure.

#0145
IMPLANTATION OF THE EDWARDS SAPIEN 3 TRANSCATHETER HEART VALVE IN REGURGITANT NON-CONDUIT RIGHT VENTRICULAR OUTFLOW TRACTS: IMMEDIATE RESULTS.
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Background: Pulmonary valve replacement (PVR) is indicated in patients with significant pulmonary regurgitation (PR). While once an exclusively surgical procedure, many patients now have PVR accomplished using transcatheter techniques with excellent early outcomes. Current literature is limited to the Melody valve and the Edwards SAPIEN XT. We aim to evaluate the procedural and immediate post-procedural results of PVR with the newest generation Edwards SAPIEN 3 valve in regurgitant non-conduit RV outflow tracts (RVOT).

Methods: After receiving IRB approval, a retrospective review of all transcatheter pulmonary valve replacements performed in our institution was conducted. We included all patients receiving SAPIEN 3 valves in regurgitant non-conduit RVOTs and excluded all others. Pre-procedural data included patient demographics, anatomy, surgical history, and baseline assessment of tricuspid insufficiency, PR and RV size/function. Procedural data included access site(s), baseline RV pressure, RVOT gradient, RVOT dimension, subsequent stented diameter, and final RVOT dimensions with assessment of any residual valve gradient or regurgitation.

Results: The SAPIEN 3 valve was implanted in 12 patients with non-conduit RVOTs between July 2015 and October 2016. Mean body weight was 61 kg (31-100). Successful valve deployment was achieved in all 12 patients (10 via transfemoral approach, 2 via Hybrid surgical approach). All patients had free PR on baseline assessment without significant RVOT stenosis. Mean RV pressure at baseline was 30 mmHg (24-37) with minimal RVOT pressure gradient (0-7). Mean RVOT dimension during balloon sizing was 28 x 27 mm. Pre-stenting was performed in 10 (77%), and 8 of those 10 underwent staged intervention. All of the 12 patients received 29 mm valves. Rapid ventricular pacing was utilized during valve deployment in all 11 patients undergoing percutaneous delivery. There were no procedural complications. Immediate results demonstrated excellent resolution of PR (trace in 10, none in 2) with minimal residual pressure gradient on pullback measurements or follow-up echocardiography. Tricuspid regurgitation was unchanged. Median length of stay was 1.5 days (1-6 days).

Conclusions: The SAPIEN 3 valve is well suited for use in regurgitant non-conduit RVOTs demonstrating excellent immediate results, but additional studies are needed to make meaningful comparisons to the historical outcomes with surgical PVR.

#0146
TRANSCATHETER CLOSURE OF MEMBRANOUS AND MUSCULAR OUTFLOW VSD’S USING A RETROGRADE ARTERIAL APPROACH.
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Methods: Peri-procedural and follow-up analysis of patients undergoing retrograde left ventricular approach. Hemodynamic instability particularly in small infants. We report our recent experience with transcatheter VSD device delivery via a direct retrograde left ventricular approach.

Results: 22 patients (16 paediatric, 9 female) with a median age 7.5 years (range 6 months to 59 years), median weight 24.3kg (range 6kg to 80kg) underwent retrograde transcatheter VSD closure. Nineteen VSDs were perimembranous, two were muscular and one case involved an LV to RA shunt. The median VSD diameter was 6mm (range 3.5mm to 11mm). Indications for closure included left ventricular volume overload (n=13), failure to thrive (n=2), previous bacterial endocarditis infection (n=2), haemodynamically significant (n=2), symptomatic (n=2) and left ventricular dysfunction (n=1). A variety of closure devices were used including Lifetech Symmetric Membranous (n=12), Amplatzer Ductal Occluder II (n=5), Lifetech Asymmetric Membranous (n=3), Amplatzer muscular (n=1), Lifetech Muscular (n=1) and Lifetech Eccentric Membranous (n=1). Mean fluoroscopy time was 15.6mins (range 5.6 to 33.2 minutes). Device delivery was successful in all cases. In one patient, the device prolapsed through the VSD during an antegrade AV loop approach. The retrograde approach was then successfully used to close the defect with a larger occluder. Immediate complete closure was seen in 15 cases with trivial to mild leak seen in the remaining 7 patients. There was no increase in the pre-procedural grading of aortic incompetence. One patient developed intermittent junctional rhythm within 24 hours however this resolved spontaneously. At median follow-up of 2.1 months (range 1.25 to 7 months), all patients were well. There were no rhythm abnormalities seen at follow-up. Trivial to mild residual leaks remained in four patients.

Conclusions: Retrograde transcatheter VSD closure is safe and effective and may avoid some of the technical challenges associated with creation of an arteriovenous loop.

#0147
CLASSIFICATION SCHEME FOR DUCTAL MORPHOLOGY IN CYANOTIC PATIENTS WITH DUCTAL DEPENDENT PULMONARY BLOOD FLOW AND INFLUENCE ON OUTCOMES OF PATENT DUCTUS ARTERIOSUS STENTING

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Introduction: Stenting of the patent ductus arteriosus (PDA) in cyanotic patients (pts) with ductal dependent pulmonary blood flow (PBF) can be technically challenging. The purpose of this study was to devise a classification scheme for ductal morphology in this population and to evaluate its association with outcomes of PDA stent implantation.

Methods: Between 2008-15 all pts < 1 yr of age who underwent PDA stenting for PBF at the 4 centers comprising the Congenital Catheterization Research Collaborative (CCRC) were included. A classification scheme for PDA morphology was devised based on tortuosity index (TI) (1-mild: relatively straight, II-moderate: 1 turn, III-severe: >1 turn); and curvature index (CI) based on the entire length of the PDA including all curves (L1) and the direct distance between aortic and pulmonary artery (PA) insertion of the PDA (L2), with CI= (L1–L2)/L1. Primary outcome was reintervention to treat cyanosis (RI) and secondary outcomes included procedure time, branch PA jailing, and subsequent surgical PA plasty.

Results: 107 pts underwent PDA stenting in whom the TI was type I (n= 61, 57%), type II (n=23, 21%) and type III (n=23, 21%). Pts with a higher CI had greater discrepancy between the pre-intervention PDA length and ultimate stented PDA length (r = 0.3, 95% CI 0.11-0.47). 46 (41%) pts required RI, which was less common in pts with TI of I versus II/III (34% vs. 50%, p=0.1). Procedure times did not differ based on TI or CI. Partial (n=18) and complete (n=4) PA jailing occurred in 22 pts. Higher TI and CI were associated with PA jailing (p < 0.01 for both). PA plasty was performed in 65 pts (61%) at time of subsequent surgical repair, with no difference in risk based upon presence or absence of a jaled PA (p=0.17). There were 14 (13.1%) complications and no mortality.

Conclusions: We devised a classification scheme based on qualitative and quantitative measures of PDA morphology that may be helpful in anticipating outcomes in cyanotic pts undergoing PDA stenting. PDA stenting is achievable even in highly tortuous PDAs, although those with higher TI had greater RI rates and those with higher TI and CI were associated with higher risk of branch PA jailing, and greater discrepancy between measured PDA length and ultimate stented PDA length. Despite this, PA plasty was not more common in pts with PA jailing.

#0148
USE OF THE SAPIEN S3 VALVE FOR TRANSCATHETER VALVE REPLACEMENT IN CONGENITAL HEART DISEASE: A MULTI-INSTITUTIONAL EXPERIENCE

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Objectives: To describe a multi-center experience and techniques used for percutaneous transcatheter pulmonary valve replacement (TCPVR) using the Edwards Sapien™ S3 Valve.

Background: Off-label use of the Sapien S3 valve can allow for large diameter transcatheter valve replacement in patients with congenital
A NEW COVERED STENT FOR COMPLEX COARCTATION AND PULMONIC VALVE LANDING ZONES.

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**Introduction:** Recent evidence is providing guidance on the need for covered stent use in lesions such as coarctation of the aorta and right ventricular outflow tract stenting. Currently the only FDA approved covered stent is the Cheatham-Platinum stent, but development in this area is urgently needed. Andratec have developed the Optimus covered stent as an alternative covered stent for similar clinical indications. This is a laser cut cobalt chromium stent with a double layer of PTFE sandwiching the stent struts. It’s hybrid design offers a wider range of expansion with less shortening and potentially more predictable behaviour of the covering. We describe the first human use of the covered Optimus stent in patients with coarctation of the aorta.

**Methods and Results:** We successfully implanted the Optimus covered stent in 7 consecutive cases in patients whose pre-procedural anatomy looked challenging for currently available covered stents (Fig 1A). Six of the patients had native coarctation with one recoarctation following surgical repair. Median age was 46yrs (range 33-69yrs). There were no significant complications with reduction in the median invasive gradient from 23mmHg to 0mmHg. The length of stents used ranged from 33mm to 57mm with a median shortening after expansion of 13%. Post-procedural follow-up with magnetic resonance imaging or computed tomography has not shown evidence of fracture or migration or re-narrowing (Fig 1B). The median duration of follow-up is 8months.

**Conclusions:** Preliminary results show that the Optimus covered stent is safe and efficacious for use in patients with coarctation of the aorta.

PERCUTANEOUS ACCESS TO ATRIAL MASS IN FOLLOWING TCPC SURGERY (MODIFIED FONTAN) FOR SINGLE VENTRICLE PHYSIOLOGY

**Methods and Results:** We successfully implanted the Optimus covered stent in 7 consecutive cases in patients whose pre-procedural anatomy looked challenging for currently available covered stents (Fig 1A). Six of the patients had native coarctation with one recoarctation following surgical repair. Median age was 46yrs (range 33-69yrs). There were no significant complications with reduction in the median invasive gradient from 23mmHg to 0mmHg. The length of stents used ranged from 33mm to 57mm with a median shortening after expansion of 13%. Post-procedural follow-up with magnetic resonance imaging or computed tomography has not shown evidence of fracture or migration or re-narrowing (Fig 1B). The median duration of follow-up is 8months.

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#0150

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#0149

A NEW COVERED STENT FOR COMPLEX COARCTATION AND PULMONIC VALVE LANDING ZONES.

Gareth Morgan, Matteo Ciuffreda, J DeGiovanni
Children's Hospital of Colorado, Denver, CO, USA

**Introduction:** Recent evidence is providing guidance on the need for covered stent use in lesions such as coarctation of the aorta and right ventricular outflow tract stenting. Currently the only FDA approved covered stent is the Cheatham-Platinum stent, but development in this area is urgently needed. Andratec have developed the Optimus covered stent as an alternative covered stent for similar clinical indications. This is a laser cut cobalt chromium stent with a double layer of PTFE sandwiching the stent struts. It’s hybrid design offers a wider range of expansion with less shortening and potentially more predictable behaviour of the covering. We describe the first human use of the covered Optimus stent in patients with coarctation of the aorta.

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**Conclusions:** Preliminary results show that the Optimus covered stent is safe and efficacious for use in patients with coarctation of the aorta.
The final palliation for single ventricle physiology these days utilise the surgical technique of Total Cavo Pulmonary Connection (TCPC) using a ptfe conduit between the inferior naval vein and the pulmonary arteries. This concept is considered to favour haemodynamics and reduce atrial arrhythmias. Whilst surgical fenestrations are sometimes performed these can get occluded; in those where there is no fenestration, entry into the LA mass through the conduit is not easy, sometimes impossible and not without risk. We describe a technique to enter the atria mass (AM) from the right internal jugular vein, mainly using a transeptal needle. The proximity of the pulmonary artery (PA) to enter the atria mass (AM) from the right internal jugular vein, mainly using a ptfe conduit between the inferior naval vein and the pulmonary arteries. This concept is considered to favour haemodynamics and reduce atrial arrhythmias. Whilst surgical fenestrations are sometimes performed these can get occluded; in those where there is no fenestration, entry into the LA mass through the conduit is not easy, sometimes impossible and not without risk. We describe a technique to enter the atria mass (AM) from the right internal jugular vein, mainly using a transeptal needle. The proximity of the pulmonary artery (PA) just to the left of the conduit anastomosis and the left atrium makes this feasible, it means crossing native tissue only and previous surgery and adhesions make the procedure safe. The procedure is carried out under general anaesthesia and with TEE guidance. A CT prior to the procedure helps with a better understanding of the anatomy although this is not mandatory. Angiography is carried out simultaneously from the superior naval vein and the left atrium (entered retrogradely via the aorta and ventricle through the AV valve). This helps with the puncture site, direction of needle and distance between the PA and AM. Pressure monitoring is important during puncture. Once the AM is entered and confirmed on pressure, TEE and angiography, the planned procedure can proceed after heparinisation. We carried out this procedure in 6 patients, 5 children and 1 adult. The ages of the children ranged from 4 to 11 years and the adult was 30 years old. In 4 of the cases, the procedure was mainly carried out to create a stent fenestration, 2 within weeks of the TCPC and 2 much later for protein losing enteropathy (PLE). In 2 patients, concomitant IVC/conduit anastomosis stenting was carried out one with a CP stent and one with an Optimus stent because of documented stenosis. One older child required access to the AM for electrophysiology and radio frequency ablation and the adult patient required a permanent transvenous atrial pacing lead because of exit block on a chronic atrial lead which was problematic to implant many years earlier. The procedures were all successful. In one patient, there was early closure of the stent fenestration and this was initially managed with i.v. TPA and once flow was reinstated, the stent was dilated with a balloon and has remained open an with no recurrence of the PLE. The stent used was an Andrastent XXL as this is what was available but these stents are designed for larger diameters. An “X” model would have been better. Access to the AM in TCPC Fontan is increasing, in part as this population is rising, they are living longer and hence will develop problems with age. Although we have used this technique safely and showed proof of concept for fenestration, permanent transvenous pacing and for arrhythmia therapy, other indications will arise in future, such as AV valve repair, left atrial appendage occlusion, pulmonary vein isolation and paravalvular leaks. Potentially, we could also see indications for percutaneous or hybrid AV valve replacement which will make this technique essential in TCPC physiology.

**Figure 1 (#0150).**

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**#0151**

**UTILIZING FLUOROSCOPY TO IDENTIFY THE POSITION OF THE ENDOTRACHEAL TUBE PRIOR TO CATHETERIZATION: A QUALITY INITIATIVE**

Daniel Gruenstein, Melissa Webb, Ala Soofian, Shari Slyder, Caitlyn Aveyard

Daniel Gruenstein, Chicago, IL, USA

**Background:** Unrecognized or delayed recognition of endotracheal tube malposition during cardiac catheterizations for congenital heart disease may lead to avoidable complications in the cath lab. Consequences include inaccurate hemodynamic data from hypoventilation, or patient complications from inadequate gas exchange. A quality initiative of fluoroscopically verifying and correcting endotracheal tube (ETT) position before obtaining hemodynamics during catheterization was implemented. We describe the frequency of ETT repositioning with this practice.

**Methods:** All patients undergoing cardiac catheterization since implementing this practice were retrospectively reviewed. Primary outcome of the study was the frequency of ETT repositioning.

**Results:** Of the 66 patients who underwent catheterization, 3 had a tracheostomy tube or ETT already in place. Of the remaining 63 patients, 17 (27%) required repositioning due to high or low placement. The mean weight and age for those requiring repositioning was 15.0 kg (2.9 – 78) and 3.3 years (0.1 – 20) and for those who did not require repositioning was 23.3 kg (3.1 – 104.8) and 5.7 years (0 – 37) respectively.

**Conclusions:** Our study demonstrates that improper ETT position is common prior to cardiac catheterization. Implementing a fluoroscopic check and correction of the ETT position prior to starting the procedure is simple, low risk, and it may improve the quality of hemodynamic data and avoid preventable complications. If broader
implementation of this practice demonstrates similar early recognition of inadequate ETT position, this simple practice may be recommended as the standard for all laboratories.

#0152
RADIAL ARTERIAL ACCESS FOR CARDIAC CATHETERIZATION: CASE REPORT
Daniel Gruenstein, Melissa Webb, S Nathan, Brojendra Agarwala
University of Chicago, Chicago, IL, USA

Background: Alternative access for percutaneous interventions is becoming more common for pediatric patients with difficult access. The smallest reported child receiving radial access for cardiac catheterization is 39 kg and youngest 12.1 years.

Methods: A 7 year old child with Tetralogy of Fallot, pulmonary atresia, and MAPCAS status post RV-PA conduit and pulmonary artery unifocalization required conduit stenting due to severe right ventricle hypertension and conduit gradient. He had bilateral femoral artery occlusion from previous cardiac catheterizations. Left radial artery access was used for left heart catheterization and conduit balloon-occlusion testing for coronary compression angiograms. Post-procedural testing showed no vascular injury a patent and intact radial artery.

Conclusion: Successful radial arterial catheterization is possible without complication in the smaller and younger patients than previously reported. This approach may provide safe opportunities complete procedures in patients with difficult access. The specific techniques and tips discussed in this case may help operators perform radial arterial catheterization for small pediatric patients.

#0153
ADVANCED TECHNIQUES FOR PERCUTANEOUS PULMONARY VALVE REPLACEMENT USING THE MELODY VALVE.
François Godart, Ali Houeijeh, Marie-Paule Guillaume, Pauline Gras, Olivia Domanski
Cardiac Hospital, Lille, France

Percutaneous pulmonary valve replacement (PPVR) with the Melody valve (Medtronic) may be challenging in patients with unfavourable RVOT. Different advanced techniques have been proposed to overcome this problem: jailing and/or Russian dolls techniques or the folded valve techniques.

From March 2015 to October 2016, 11 patients (5F/6M) with a mean age of 19 ± 10 years (11-45 years) underwent these procedures. Initial pathology included tetralogy of Fallot (n=7), transposition of the great vessels (n=2), pulmonary valve stenosis (n=1), and aortic valve stenosis (n=1). These patients had undergone a median of 2 previous surgical repairs. The RVOT had been previously repaired with a transannular patch and 1 patient had homograft. The indications for pulmonary valve replacement were: significant pulmonary regurgitation (n=7) and a mixed lesion (n=4). All patients had before the procedure, MRI study and CT scan to delineate the exact morphology of the RVOT. Before implantation, balloon dilatation of the RVOT with control aortography to obviate any coronary artery compression was performed in all. Pre-stenting was realized in all with LD max stent (Ev3).

A 22-mm Melody valve was implanted in 9 patients, a 20-mm valve in 2 (9/11 under left ventricular pacing). The folding techniques were employed in 8 patients, the PA branch jailing in 7, and the Russian dolls technique in 3. The folding technique on both extremities of the stent (n=1) and only on distal end (n=7) was performed because of short pulmonary artery trunk with early PA bifurcation. These techniques were combined in 6 patients. RVOT dilated up to 25-26 mm in diameter could be thus corrected by PPVR with a 22 mm Melody using these techniques.

During follow-up (1 to 19 months), no patient had reintervention. No endocarditis was observed.

These advanced techniques using the Melody valve can extend the classical indications for PPVR beyond the revalvulation of conduit. Patched RVOT up to 25-26 mm in diameter can be repaired. These initial results are promising but more experience and longer follow-up are mandatory.

#0154
IS THE NEW OCCLUTECH PDA OCCLUDER A GOOD ALTERNATIVE FOR TRANSCATHETER CLOSURE OF PDA?
François Godart, Ali Houeijeh
Pediatric Cardiology, Lille, France

Purpose: One tertiary centre experience with the new Occlutech PDA occluder for arterial duct occlusion.

Methods: From March 2013 to October 2016, 29 patients (20 females and 9 males) underwent percutaneous arterial duct closure with the new Occlutech PDA occluder. All patients had significant L-to-R shunt with enlarged left ventricle.

At implantation, the median age was 26 months and median weight was 12kg (4.1 to 57 kg). The procedure was realized under local anaesthesia in the majority of patients. Size of the duct was 2.94 ± 0.85 mm on angiography. According to PDA Krichenko classification, ducts were: type A (n=21), type E (n=7) and B (n=1). The systolic pulmonary artery pressure was 42 ± 15 mm Hg. Implantation succeeded in all. Closure was performed by the standard 4/6 mm occluder (n=14), the standard 5/7 mm occluder (n=9), the standard 6/8 mm occluder (n=4), the standard 8/10 mm occluder (n=1), and the standard 3.5/5 mm (n=1) using a 6 or 7 F delivery sheath. Technique of implantation is similar to that of the Amplatzer Duct Occluder and there is no learning curve with the use of this device. After implantation, trivial shunt was noticed on angiography in 21 patients, 8 had no shunt. The mean radiation dose was 7.2 Gycm2. Neither embolization nor haemolysis was observed. After closure, femoral thrombosis was noticed in 3 patients but resolved completely under heparin therapy. On control Doppler echocardiography, 6 patients had tiny residual shunt after implantation. At the control one month later, shunt was closed in 5 of them; in the remaining one, full occlusion was observed 18 months
after implantation. No obstruction of the left pulmonary artery or isthmic stenosis was noticed. These results are very similar to those published with the Occlutech device including 171 patients from 4 articles: successful implantation (95.4–100%), full occlusion during follow-up (96-100%); complications were observed in 4 patients: femoral thrombosis (n=2) and embolization (n=2).

Conclusions: Transcatheter closure of PDA with the new Occlutech PDA occluder is a safe and effective. In fact, the results are very similar to that reported with the Amplatzer Duct Occluder. The Occlutech PDA occluder appears as a valuable therapeutic option for transcatheter occlusion of the arterial duct. Further studies with longer follow-up are necessary to confirm these results.

#0155 SUCCESSFUL TRANSCATHETER CLOSURE OF UNROOFED CORONARY SINUS USING COVERED STENTS IN AN ADULT WITH DRAINAGE OF THE CORONARY SINUS TO THE RIGHT VENTRICLE

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2Our Lady’s Children’s Hospital, Dublin, Ireland
3Rockford Memorial Hospital, Rockford, IL, USA
4Weill Cornell Medicine & Sidra Medical and Research Center, Doha, Qatar

We present a rare case of a 41 year-old female with unroofed coronary sinus who underwent supra-annular tricuspid valve replacement with consequent drainage of the coronary sinus to the right ventricle. It is unclear whether the coronary sinus was unroofed congenitally or iatrogenically (as a result of increased pressure load on the coronary sinus). Either way this rare setup resulted in a significant central cyanosis as a result of shunting the desaturated blood from the right ventricle to the left atrium. In addition, she was at high risk of developing paradoxical embolism and other complications of chronic cyanosis. This connection was successfully closed via trans-catheter delivery of covered stents and lead to resolution of the cyanosis. Post stenting transesophageal echocardiogram with contrast injection in the coronary sinus revealed no residual communication with the left atrium. Our patient recovered well from the procedure. On follow-up, the patient has normal systemic oxygen saturation with stable cardiac function and improved exercise tolerance.

Catheter-based management of unroofed coronary sinus is still evolving; there is only one case reporting deployment of a covered stent and limited cases reporting successful closure using Amplatzer® Ductal Oclcluders. Use of covered stents may be preferable as it restores normal physiologic circulation and eliminates right to left shunting. However, it may be challenging especially in complete unroofing. Furthermore, it may be more difficult to ensure stent stability and requires extreme caution to avoid obstructing coronary venous drainage into the coronary sinus. This might result in abnormalities in electrophysiological conduction.

In conclusion, transcatheter occlusion of unroofed coronary sinus, although it is technically difficult, is feasible and provides an alternative to cardiac surgery as a first line in managing unroofed coronary sinus. However, extreme caution and experienced interventionists are prerequisites to consider this type of procedure due to technical difficulties and possible complications.

#0156 RESULTS OF LD ADVANTA™ V12 COVERED STENT TRIAL FOR COARCTATION OF THE AORTA.

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Objective: To assess the safety and effectiveness of the Large Diameter Advanta™ V12 Covered Stent for treatment of coarctation of the aorta.

Methods: A prospective, multicenter, non-randomized, single arm study. Primary Efficacy Endpoint: Significant reduction in gradient assessed by echo-Doppler (diastolic velocity) and DV/SV (diastolic velocity/systolic velocity) at pre-stent vs. immediately post stent and pre-stent vs. 12 month follow-up. Primary Safety Endpoint; MAE and MAVE occurring within 30 days.

Results: Between 2009 and 2013, 70 patients [43M, 27F] of median age 17.0 yrs [6.0-75.0] and weight 57.4Kg [28.2-120] with coarctation of the aorta were enrolled. Stents were successfully implanted and in all patients. Mean CoA diameter increased from 5.5±3.6 to 13.1±3.2mm [p<0.001]. Mean peak gradient decreased from 35.8±16.0 to 5.4±7.8mmHg [p<0.001]. A Baseline DV/SV was 0.59± 0.17 dropped to 0.34 ± 0.13 [p<0.001] after stenting and was unchanged at 12 month follow-up. DV was 216.3 ± 77.5cm/s dropped to 90.9 ± 46.5cm/s [p<0.001] and was unchanged at 12 months. 61 patients underwent CTA at ~12 months with mean stent diameter maintained at 13.3 ± 2.1mm, 91 ± 0.1% of diameter of the transverse arch.

Complications: Acute MAVE was 1.4% [1 femoral artery occlusion]. MAVE at 12 months was 4.3% including 1 aortic hematoma treated with a second stent, 1 small aneurysm at distal edge of the stent treated with a second stent. At 12 months there was no new wall injury on CT. 3 patients had late (>1 yr) stent infolding requiring re-intervention.

Conclusions: The Large Diameter Advanta™ V12 Covered Stent is safe and effective for treatment in coarctation of the aorta. Long term follow-up to assess aortic wall injury and stent integrity is required.

#0157 PREDICTORS OF PROCEDURAL FAILURE IN PATIENTS UNDERGOING PERCUTANEOUS CLOSURE OF LARGE ATRIAL SEPTAL DEFECTS USING THE AMPLATZER SEPTAL OCCLUDER.

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Aims: Owing to their more complex anatomy, large atrial septal defects can present unique challenges for percutaneous closure. Previous reports observed a much higher procedural failure rate in
this specific subgroup when compared to small and moderate size defects. We sought to determine clinical and anatomical predictors of procedural failure in patients undergoing percutaneous closure of atrial septal defects larger than 24mm using the AmplatzerTM Occluder device (St.Jude Medical Inc.).

**Method & Results:** Adult patients referred between January 2006 and July 2013 for percutaneous closure of a secundum atrial septal defect (ASD) with a stretched diameter higher than 24 mm were included in this study. Clinical, imaging and procedural data were retrospectively collected from index admission to last follow-up. Patients underwent ASD closure according to standard indications under transseousphageal and fluoroscopic guidance. A total of 90 patients, mean age 47±16 years, 60% female, met inclusion criteria. Patients presented with right heart failure symptoms in 81.1% of cases, 93.3% had right ventricular enlargement and 7.8% had a prior embolic event. Median non-stretched and stretched ASD diameters were 24.0 mm (20;28) and 29.0 mm (25;34,0) respectively. Procedural success rate was 81.1%, one patient (2.2%) required emergent surgery because of a ruptured septum after sizing-balloon inflation resulting in cardiac tamponade. During a median follow-up of 19.5 months, we observed 4 (4.4%) episodes of systemic thrombo-embolism, 5 (5.5%) new onset supraventricular tachyarrhythmias, no cardiac erosion, no device embolization and no cardiovascular death.

In univariate logistic regression analysis predictors of failure were non-stretched diameter (OR=1.25 per mm; p=0.00002), stretched diameter (OR=1.6 per mm; p<0.0001), absence of any type of margin (OR=4.8, p<0.0001) and absence of postero-inferior margin (OR=39.3; p<0.0001). Multivariate regression analysis identified non-stretched diameter (OR=1.17 per mm; p=0.05), absence of any type of margin (OR=5.25; p=0.0004) and absence of postero-inferior margin (OR=100, p=0.05) as independent predictors of failure.

**Conclusion:** Success rate of transcatheter closure of large atrial septal defects using the Amplatzer device is lower when compared to small and moderate defects. A large unstretched diameter, absence of any type of margin and more strongly absence of postero-inferior margin were independently associated with procedural failure. These findings can be assessed prior to the procedure and could improve patient selection.