Introduction
Scimitar syndrome is a rare and complex association of congenital cardiopulmonary abnormalities with an estimated prevalence of 1 to 3 per 100,000 births [1]. Literature demonstrated that patients presenting in infancy are more likely to have severe heart failure, Aorto-pulmonary Collaterals, and a poorer prognosis than patients presenting at a later age [2-4].

Case History
We are presenting a 4 weeks old neonate with weight of 2.7 kg, diagnosed as a non-obstructive scimitar syndrome (right lower pulmonary vein draining directly to IVC) and multiple massive aorto-pulmonary collaterals to the right lung and ASD secundum, hypoplasia of right pulmonary artery and right lung. Diagnosis was confirmed by transthoracic Echocardiography, CT scan of abdomen and thorax with contrast and cardiac MRI.

Deploying MVP to occlude AP collateral
An Angiograph showing a deployed Piccolo Amplatzer occluder in one of the AP collaterals, and adjusting a microvascular plug in another AP collateral.

Case History
Patient was not diagnosed antenatally. Presented after birth with severe pulmonary hypertension requiring high setting of ventilation and high serum lactate level. Patient developed systemic circulation steel with bowel ischemia and infection (necrotizing enterocolitis). After failure of aggressive supportive medical treatment, patient was taken to the catheterization lab for diagnostic and interventional catheter.

Major Aorto-Pulmonary Collaterals

Descending Aorta Angiography demonstrating multiple Major Aorto-Pulmonary Collaterals arising from Abdominal Aorta to the right lung
Deploying multiple occluders for AP collaterals

An angiograph showing multiple occluders deployed in the AP collaterals

An angiograph showing multiple occluders deployed in the AP collaterals, with remaining collateral that is occluded in the last angiograph

Therapeutic Interventions
Angiographies showed multiple massive collaterals from abdominal aorta to the right lung with pulmonary venous return of the lower right lobe of the lung to the IVC and significant cardiomegaly.

We used multiple microvascular plugs, PDA occlusion devices, Piccolo device through 3 Fr then 4 Fr sheaths and catheter through right femoral artery to occlude the massive collaterals.

Outcome
Post-procedure patient was extubated in the NICU and heart failure symptoms disappeared.

Conclusions
Massive aortopulmonary collaterals transcatheter arterial embolization can be done safely in the catheterization lab in neonates with small body weight and represent a minimal invasive option compared to surgical option. Embolization of anomalous systemic arteries can dramatically improve symptoms of heart failure and pulmonary hypertension in neonates.

References