30% within the not negative groups (excluding SADS where it was 100%). Factors contributing to low rates of genetic testing and phenotypic predictors of positive genotype warrant further investigation.

Moderated posters

23 REAL-LIFE USE OF IMPELLA® CP: THE NORTHERN IRISH EXPERIENCE
U Azhar Khan, M Saad Ahmed, J Conleth Murphy. Royal Victoria Hospital, Belfast, UK
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Introduction The Impella® CP (ABIOMED Inc., Danvers MA, USA) is a relatively novel percutaneously inserted mechanical circulatory support (MCS) system. It is a 14 Fr micro-axial temporary left ventricular assist device, which provides continuous forward flow of up to 4.3 L/min into the ascending aorta to augment cardiac output. It is indicated to provide additional haemodynamic support in cases of advanced cardiogenic shock (CS), primarily in acute myocardial infarction (AMI). It is also used in selected high-risk patients undergoing complex percutaneous coronary intervention (PCI). It is a widely known modality but data from randomized controlled trials (RCT’s) have not demonstrated a clear clinical benefit. We aim to describe our experience with use of Impella® CP in our institution.

Methods The Belfast Health & Social Care Trust is a high-volume interventional centre, which performs approximately 2500-3000 PCI per annum, of which 650 are emergency procedures. We retrospectively reviewed all patients who were treated with Impella® CP for cardiogenic shock after acute myocardial infarction as well as those undergoing elective high-risk PCI [table 1]. We report patient demographics, device-related complications and clinical outcomes in these cohorts.

Results Since March 2018, 7 patients have received Impella® CP support. 5 patients presented with AMI complicated by CS, and the remainder 2 had elective high-risk PCI. Mean age in the CS cohort was 49.8 years, with an average of one case every 232 days. Mean age in the elective high-risk PCI cohort was 73 years. In the CS cohort, 3 patients were subsequently transferred to a quaternary transplant centre with subsequent advanced heart failure (HF) input (2 extracorporeal membrane oxygenation and 1 surgical left ventricular assist device). 1 CS patient was transferred but did not undergo advanced HF input and was palliated. 1 CS patient died in the cathlab. In the total cohort 4/7 (56%) of patients had direct device-related complications (2/7 limb ischaemia, 2/7 bleeding requiring blood transfusions). Overall 30-day mortality in the CS cohort was 40% (2/5).

Conclusion Impella® CP use in our high-volume center is infrequent and carries a high complication rate. Overall mortality rate in our CS group remains high but it is difficult to make definitive conclusions on small numbers. Our complication rate is significant, but similar to previous documented series in higher-volume centres. Impella® CP may have clinical benefits in carefully selected patients, but in the absence of data from robust multi-centre RCT’s patient selection will remain speculative. The exact role for Impella® CP remains unclear.

24 SHOCKWAVE INTRAVASCULAR LITHOTRIPSY AS AN ADJUNCT TO PERCUTANEOUS CORONARY INTERVENTION IN CALCIFIED CORONARY LESIONS: A 2 YEAR, SINGLE CENTRE EXPERIENCE
H Muragagha, K Millar, B Wei Soh, I Ullah, T Hennessy, S Amous, T Kiernan. University Hospital Limerick, Ireland
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Background Calcified coronary lesions are attributed to multiple risk factors such as increasing age, smoking, DM, and CKD. PCI of calcified lesions is challenging due to impaired stent crossing, under-expansion, and mal-apposition. Shockwave intravascular lithotripsy (S-IVL) is a novel angioplasty balloon that emits sonic pulsations, thus disrupting intra-coronary calcium and often obviates the need for conventional rotational atherectomy (RA). There is limited real-world data available on the use of S-IVL and we aim to review its use in our center.

Methods A retrospective single-center cohort study was performed of all S-IVL cases between Sep 2018 and Apr 2021.

Results Thirty-seven S-IVL procedures were performed during the period of the study. The mean age is 72.32 years (range 65-87, Median age 74.76%) were male. All patients had moderate to severely calcified coronary artery lesions. Indications for PCI included: 24 unstable angina, 9 NSTEMI, 3 Stable CAD, 1 ST-elevation MI, and 2 recurrent ISR despite drug-eluting balloon therapy Risk factors included: 11 (29.7%) DM, 34 (91.8%) HTN, 32 (86.4%) dyslipidaemia, 9 (24.3%) active smoking, 7 (18.9%) CKD, 2 (5.4%) previous TIA, 3 (8.1%) previous CABG and 13 (35.1%) previous PCI. All lesions were post-dilated, mainly by using a 3.5 mm diameter; high pressure was also applied in almost 90% of post-dilation balloons. IVUS was used in 5 cases (mainly in LM procedure) to