Purpose To investigate differences in plasma clot properties in patients with AF and CAD and compare the effect of warfarin and antiplatelets on clot structure in AF population.

Methods We studied 270 patients and divided them into 3 groups: AF on warfarin (n=184), AF on antiplatelets (n=46) and CAD (n=40). Plasma samples were obtained from participants and centrifuged to prepare platelet poor plasma. Assays were performed in 96-well polystyrene microtiter plates. Reagents were diluted in standard buffer (10 mM N-2-hydroxyethylpiperazine-N’-2-ethanesulphonic acid [HEPES], pH 7.4, 150 mM NaCl). Patient plasma samples (25 %) were incubated with tissue plasminogen activator (500 ng.ml-1) for 10 minutes at 37°C before the addition of CaCl2 (7.5 mM). Either PPP reagent (2.5 %), aPPT reagent (2.5 %), or thrombin (0.5 U.ml-1) were then added to initiate coagulation. Polymerisation of fibrin in plasma was monitored (ΔOD340 nm) using a Synergy H1 hybrid multi-mode plate reader, readings were taken in 12 second intervals for up to 60 minutes.

Results Comparisons between the 3 groups was performed using Kruskal-Wallis test, with Dunn’s post-hoc analysis and Holm-Sidak adjustment. There were no significant differences in clot structure between 3 subgroups. The maximum rate of clot formation was significantly delayed in the warfarin subgroup with all reagents used (p<0.001) (table 1). Plasma clot susceptibility to fibrinolysis increased with warfarin compared to antiplatelets but was significant only with APPT and thrombin reagents (p<0.001 and 0.04 respectively).

Conclusion Warfarin was effective in delaying clot formation compared to antiplatelets and also resulted in increased susceptibility of plasma clot to fibrinolysis.

Conflict of interest None

**BS43** _ABSTRACT WITHDRAWN_

**BS44** _CYTOKINE INDUCED DOWNREGULATION OF Plasma Membrane Calcium ATPase 4 GENE INCREASES SENSITIVITY TO APOPTOSIS IN PULMONARY ARTERY ENDOTHELIAL CELLS_

1Jude Ihuqb*a, 2Satishkumar Kunarsamy, 2Reshma Naomi Ranjit Immanuel, 2Kiruba Khan, 3Jayashree Jayachandran, 4Nadine Arnold, 4Priscille Polla, 4Fabio Gomez-del Arco, 4Juan Miguel Redondo, 4James Cotton, 5Paul D Upton, 5Nicholas Morrell, 6Allan Lawrie, 2Angel L Armesilla. 1Research Institute in Healthcare Science; 2RIHS, FSE, University of Wolverhampton; 3Department of Infection, Immunity and Cardiovascular Disease, University of Sheffield, Sheffield, UK; 4IER, Instituto de Salud Carlos III, Madrid, Spain; 5Centro Nacional de Investigaciones Cardiovasculares, Madrid, Spain; 6Department of Cardiology, Heart and Lung Centre, New Cross Hospital, Wolverhampton, UK; 7Department of Medicine, University of Cambridge, Addenbrooke’s and Papworth Hospitals, Cambridge, UK; 8University of Cambridge

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Introduction Pulmonary arterial hypertension (PAH) is a life-threatening disease characterized by progressive vasoconstriction, vascular remodelling, and occlusion of small pulmonary arteries. It leads to increased pulmonary resistance and finally right ventricular failure. The disorder has no cure and current therapies only target vasoconstriction but have little effect on vessel remodelling. Increased activity of pro-inflammatory cytokines is linked to PAH pathogenesis. In this study, we analysed the effect of TNF-alpha and IL-1Beta on the expression of Plasma Membrane Calcium ATPase 4 (PMCA4) in pulmonary artery endothelial cells (PAEC).

Methods PAEC were cultured for different times and with different doses of TNF-alpha or IL-1Beta. Expression of PMCA4 RNA and protein was determined by qPCR and western blot respectively. PMCA4 expression was silenced using siRNA specific for human PMCA4. Quantification of apoptotic cells was performed by flow cytometry and TUNNEL.

Results Treatment of PAEC with TNF-alpha or IL-1Beta induced a time- and dose-dependent decrease in the levels of RNA for PMCA4. Analysis of PMCA4 RNA levels in the lungs of mice with overexpression of ectopic TNF-alpha confirmed the in vivo relevance of these observations. RNA decay experiments performed by blocking cellular transcription with Actinomycin D indicate that the downregulation of PMCA4 RNA levels mediated by pro-inflammatory stimuli in PAEC is the result of a decrease in RNA stability. In agreement with the reduction observed in RNA levels, PMCA4 protein expression was strongly decreased by treating PAEC with TNF-alpha or IL-1Beta. Silencing PMCA4 gene expression sensitised PAEC to apoptosis, suggesting that PMCA4 protects PAEC to apoptosis induced by pro-inflammatory cytokines.

Conclusion The pro-inflammatory cytokines TNF-alpha and IL-1Beta significantly downregulate the expression of the PMCA4 gene in PAEC at the RNA and protein level. Decrease in PMCA4 expression sensitised PAEC to apoptosis. This indicates that the PMCA 4 gene might play an important role in the apoptotic loss of endothelial cells observed in the pulmonary arterioles of patients with PAH.

Conflict of interest None

**BS45** _ACTIVATING TRANSCRIPTION FACTOR ATF2 NEGATIVELY REGULATES THE EXPRESSION OF ENDOTHELIAL NOTCH LIGANDS_

1Ivonne Olivaesa, 2Kirthika Kalyanakrishnan, 2Suhaib Ahmed, 2Clare Muccott, 2Robert N Wilkinson, 3James Cotton, 4Wolfgang Breitwieser, 5Mark R Morris, 1Angel L Armesilla. 1RIHS, FSE, University of Wolverhampton; 2Department of Infection, Immunity and Cardiovascular Disease & Bateman Centre, University of Sheffield; 3Department of Cardiology, Heart and Lung Centre, New Cross Hospital, Wolverhampton, UK; 4Molecular Biology Core Facility, Cancer Research UK, Manchester, UK

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Introduction ATF2 also known as cyclic AMP response element binding protein 2 (CREB2) is a member of the leucine zipper (bZIP) transcription factor family that binds to specific DNA sequences and regulates the transcriptional activation of target genes. ATF2 regulation and functions have been studied in a number of developmental and pathological conditions. Studies have shown that activation of ATF2 by VEGF mediates angiogenic processes such as endothelial cell migration and tubular morphogenesis but the molecular role of ATF2 in these processes is largely unknown. To shed some light on this matter we are identifying ATF2-target genes induced by stimulation of endothelial cells with pro-angiogenic stimuli.

Methods HUVEC were stimulated with various pro-angiogenic stimuli at different times, and the activation (phosphorylation) status of ATF2 was determined by Western Blot.

ATF2 functionality in endothelial cells was suppressed by infecting HUVEC with an adenovirus encoding a phosphorylation-mutant, dominant-negative version of ATF2 (Ad-ATF2AA) where phosphorylation residues Thr69 and Thr71 have been mutated to Ala. HUVEC infection with an adenovirus encoding GFP was used as a control. PCR-based screening of specific gene arrays was used to identify the effect of ATF2 loss-