and fibrosis in myocardium, thus delay the progress of the diabetic cardiomyopathy.

**e0041** **VASOMOTOR FUNCTION FOLLOWING NEWER GENERATION OF BARE METAL STENT OVERSTRETCH IN A PORCINE CORONARY MODEL**

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**Backgrounds** Overstretch damage after bare metal stent (BMS) placement could trigger cell proliferation and in-stent restenosis (ISR). Newer Co-Cr BMS has thinner stent struts, which designs to minimise cellular response to injury. We aimed to investigate neointimal growth, as well as vasomotor function after overstretch using Co-Cr BMS in a pig coronary model.

**Methods** 15 vessels in five pigs were assigned to receive BMS (stent struts 91 μm) implantation with either S/A ratio 1.3 (group I, n=7) or 1.5 (group II, n=8). Quantitative coronary angiography (QCA) and optical coherence tomography (OCT) were performed at 14 days after stent implantation. Coronary vasomotor function was evaluated by incremental acetylcholine (Ach) (10⁻¹⁰ and 10⁻⁶ M) and nitroglycerin (NTG, 400 μg) infusion before stent implantation and at 14 days. Endothelial response to Ach was measured at 5–10 mm distal to the stent edge.

**Results** Both QCA and OCT showed that in-stent stenosis of group I were significantly smaller than group II at 14 days (QCA-late loss (LL), 1.22±0.21 mm vs 1.79±0.17 mm; OCT % AS, 17.0±7.9 vs 26.9±10.7% at 14 days, p<0.05 and 0.001, respectively). Liner regression analysis QCA-LL is proportional to obtained S/A ratio (r=0.60, p<0.05). Endothelium-dependent vasomotion at distal non-stented reference segments was no difference between groups. The mean coronary diameter changes at Ach 10⁻⁷ M and 10⁻⁶ M was 2.1±0.2% and 2.1±0.2% in group I; 2.2±0.2% and 2.1±0.2% in group II (p>0.05, accordingly). There was also no difference before and at 14 days after stent implantation.

**Conclusion** The progression of neointimal hyperplasia after BMS implantation is positively associated with the extent of coronary artery injury. Coronary endothelial function is preserved after BMS implantation at 14 days, which is independently of overstretch degree.

**e0042** **THE EFFECT OF ISCHAEMIC POSTCONDITIONING ON THE STRUCTURE, FUNCTION AND CX43 OF MITOCHONDRIA IN RABBIT MYOCARDIAL ISCHEMIA/REPERFUSION INJURY**

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**Objective** To investigate the effects of ischaemic postconditioning on structural and functional and connexin 43(Cx43) changes of mitochondria induced by myocardial ischaemia reperfusion (IR) injury of rabbits in vivo and potential mechanism.

**Methods** In anaesthetized open-chest rabbits, the left anterior descending artery (LAD) was occluded for 30 min and reperfused for 4 h. Sixty-four rabbits were randomly divided into four groups (n=16 in each group): Sham operation group (Group Sham), ischaemic reperfusion group (Group IR), ischaemic preconditioning group (Group IP) and ischaemic postconditioning group (Group FC) with sixteen rabbits in each. All rabbits in the four groups were killed 4 h after reperfusion. Myocardial infarct size were determined at the end of the experiment. Mitochondria were isolated with different centrifugations. Ultrastructural changes of mitochondria were observed under electronmicroscope and mitochondrial membrane potential, Ca²⁺ concentration, MDA content and SOD activity of myocardial mitochondria were examined. The content of the mitochondria Cx43 were detected with Western Blot.

**Results** Myocardial infarct size was significantly reduced in IP (18.9±2.8%)and FC (19.1±5.9%) groups as compared to IR groups (35.7±5.8%), p<0.01). Compared with group IR, the damage of mitochondrial ultrastructure were milder and Ca²⁺ concentration and MDA content were much lower in group IP and group FC (p<0.05). Mitochondrial membrane potential (p<0.01) and SOD activity of myocardial mitochondria in group IP and group FC was significantly higher than that in group IR (p<0.05). Compared with sham group, the mitochondria Cx43 expression is distinctly decreased compared group IR (p<0.05) and no significant difference was found between Group IP and Group FC.

**Conclusion** FC can protect mitochondrial ultrastructure by increasing mitochondrial membrane potential and SOD activity, and by alleviating Ca²⁺ overload, and by decreasing MDA content in myocardial mitochondria. The mechanism of FC protection to mitochondria may be concerned with FC attenuating the decrease the mitochondria Cx43 expression induced by ischaemia/reperfusion injury.

**e0043** **EFFECTS OF SIMVASTATIN ON ANGOGENESIS AND THE EXPRESSION OF ANG1 AFTER MYOCARDIAL INFARCTION IN RATS**

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**Objective** To investigate the effects of simvastatin on myocardial angiogenesis and the expression of angioptoin-1 after experimental myocardial infarction (MI) in rats.

**Methods** 60 healthy adult SD rats were randomly divided into the sham operated group, the control group, high dose of simvastatin (1 mg·kg⁻¹·d⁻¹) group, medium dose of simvastatin (10 mg·kg⁻¹·d⁻¹) group, high dose of simvastatin (10 mg·kg⁻¹·d⁻¹) group, low dose of simvastatin (40 mg·kg⁻¹·d⁻¹) group, medium dose of simvastatin (10 mg·kg⁻¹·d⁻¹) group, and control group. Rats were administered simvastatin respectively via oral gavage for four consecutive weeks starting at the next day. Density of new microvessels in the ischaemic area, LVMI, protein and mRNA expression of Ang-1 were detected 4 weeks after operation.

**Results** (1) Compared with the control group, the Density of new microvessels in low and medium dose of simvastatin group increased significantly (p<0.05); and those did not changed significantly in high dose of simvastatin group (p>0.05). (2) LVMI in low and medium dose of simvastatin group decreased significantly compared with that in control group (p<0.05), and further decreased in high dose of simvastatin group. (3) The protein and mRNA expression of Ang-1 in all simvastatin group increased significantly compared with that in control group (p<0.05).

**Conclusion** (1) Low and medium dose of simvastatin can stimulate myocardial angiogenesis after MI, whereas high dose of simvastatin have no pro-angiogenic effect. (2) The pro-angiogenic effect of simvastatin may be associated with upregulated expression of Ang-1.

**e0044** **THE ROLE OF ANG1 AND ENOS IN THE PRO-ANGIOGENIC EFFECT OF SIMVASTATIN AFTER MYOCARDIAL INFARCTION IN RATS**

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**Objective** To investigate the roles of angioptoinet-1 (Ang-1) and endothelial nitric oxide synthase (eNOS) in pro-angiogenic...