**Results** 1. In addition to expressing characteristic hHCN4 protein, mHCN4-transfected hMSCs also express an anticipated high level of hHCN4 gene by RT-PCR and Western blot analysis. And immunofluorescence image is shown for GFP. Control MSCs were negative. 2. With the use of the whole cell configuration of the patch-clamp technique, I_T was elicited using hyperpolarizing steps in 10-mV increments from -40 mV to -140 mV from a holding potential of -40 mV and was voltage-dependent. The threshold of voltage for activation of I_T is around -80 mV. Remarkably, all hHCN4 positive cells exhibit a large caesium-sensitive I_T and it was significantly inhibited by 4 mM caesium chloride.

**Conclusions** The pacemaker current of I_T can be elicited from the mesenchymal stem cells transfected with HCN4 genes by Lentiv. The genetically engineered MSC expressing hHCN4 is a demonstration of feasibility of preparing MSC-based biological pacemaker cells.

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**e0032 ROLE OF ATRIAL SUBSTRATE REMODELLING IN INDUCIBILITY OF ATRIAL FIBRILLATION AFTER EPICARDIAL GANGLIONIC PLEXI ABLATION**

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**Objective** Investigating the long-term effect of ganglionated plexi (GP) ablation on atrial fibrillation (AF) after GP ablation.

**Methods** 13 dogs were randomly divided into sham-operated group and GP ablation group. All animals underwent a right thoracotomy at the 4th intercostal space. Induced AF and atrial effective refractory period (AERP) were measured by burst rapid pacing at right atrium. After anterior right GP and inferior right GP ablation, AF and AERP were measured again in the GP ablation group. The animals were allowed to recover for 8 weeks, after which, AF and AERP was measured again. The levels of atrial natriuretic peptide (ANP), TNF-a and interleukin (IL)-6 in blood and atrial tissues were measured again. The levels of atrial natriuretic peptide (ANP), TNF-a and interleukin (IL)-6 in blood and atrial tissues were measured.

**Results** AF was easily induced in the GP ablation group after 8 weeks while AF was not observed in the sham-operated group, and immediately after GP ablation. AERP and dispersion of AERP (dAERP) were increased after GP ablation, while AERP recovered after 8 weeks. Compared with sham-operated group, the levels of ANP, TNF-a and IL-6 in the right atrium increased significantly 8 weeks after GP ablation. (204.6±31.2 vs 299.1±52.5; 1.3±0.5 vs 4.7±0.7; 0.9±0.3 vs 1.8±0.5; p < 0.05). In GP ablation group, the density of GAP43-positive, TH-positive and ChAT-positive nerves in the right atrium was 821±591; 481±627 and 629±644 per mm², respectively, which was significantly (p <0.01) lower than the nerve density in sham-operated tissues (2590±841, 1752±605 and 3147±386 per mm², respectively).

**Conclusion** Atrial substrate remodelling after GP ablation may be the mechanism of induced AF.

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**e0033 MECHANISMS OF LOSARTAN ATTENUATES VENTRICULAR REMODELLING AND CARDIAC FUNCTION IN AORTIC BANDED RATS WITH CHRONIC HEART FAILURE**

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**Objective** To investigate the regulation of Losartan (Los) on ventricular remodelling and cardiac functioning chronic heart failure rats induced by aortic banded.

**Methods** Sprague-Dawley (SD) rats underwent abdominal aorta coarctation to induce CHF confirmed by ultrasound cardiograph and Catheterisation, or sham operation, followed by 8 weeks treatment with Losartan (20 mg/kg per day, orally) or vehicle (drinking water). Plasma norepinephrine (NE) was measured by ELISA, and plasma and tissue angiotensin II (Ang II) levels were measured using RIA. Cardiomyocyte apoptosis was examined by agarose gel electrophoresis and TUNEL's method. The mRNA levels of Bax and Bcl-2 were determined by RT-PCR and the protein expression of phosphorylated and total Akt (p-Akt) and p-Akt) were assessed by Western blot.

**Results** Losartan-treated CHF rats had lower left ventricular end-diastolic pressure (LVEDP) [COA+Los: (11.47±5.06) mm Hg vs COA+Vehicle: (21.18±6.56) mm Hg, p<0.01], higher left ventricular ejection fraction (LVEF) [COA+Los: (63.28±4.32)% vs COA+Vehicle: (45.27±4.25)%], p<0.05 and lower plasma NE[COA+Los: (2.41±0.15) mg/g vs COA+Vehicle: (3.12±0.15) mg/g, p<0.05], lower plasma NE[COA+Los: (62.24±6.46) pg/ml vs COA+Vehicle : (908.24±75.10) pg/ml, p<0.05] and myocardium Ang II[COA+Los : (60.15±6.22) pg/mgprot vs COA+Vehicle : (92.31±3.1) pg/mgprot, p<0.05], but higher plasma Ang II[COA+Los : (629.68±58.71) pg/ml vs COA+Vehicle : (478.67±51.12) pg/ml, p<0.05] than vehicle-treated CHF rats. Losartan-treated HF rats had no obviously “DNA ladder” which was the character of apoptosis, and the apoptosis index was also reduced [COA+Los: (5.49±0.45)% vs COA+Vehicle: (9.62±0.51)%], p<0.05 with a lower express of Bax/Bcl-2 gene [COA+Los: (109.58±7.53)% vs COA+Vehicle: (136.76±8.82)%], p<0.05 and a higher protein expression of p-Akt[COA+Los: (70.80±6.40)% vs COA+Vehicle: (50.30±4.80)%, p<0.05].

**Conclusion** Losartan might inhibit cardiomyocyte apoptosis and improve cardiac function in aortic banded rats by blocking Ang II to bind AT1-R and promoting the activation of Akt.

**e0034 INHIBITION OF ATORVASTATIN ON THE AUTOPHAGY OF VASCULAR ENDOTHELIAL CELLS**

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**Objective** To explore the mechanism of atorvastatin’s protection on vascular endothelial cells, we conducted the research of impact of atorvastatin on vascular endothelial cells autophagy in different times.

**Methods** We used the Hank’s to replace the normal medium to induce autophagy of vascular endothelial cells. In pre-induction and induction procedure, the cells were incubated with normal medium which includes atorvastatin or not and Hank’s, respectively. Experimental cells were randomly divided into four groups: control group (Group I), the group of pre-induction with atorvastatin (Group II), the group of induction with atorvastatin (Group III) and the group of pre-induction and induction both with atorvastatin (Group IV). Transmission electron microscope (TEM) was used to detect autophagy. The RT-PCR was employed to detect the autophagy-specific markers (Bclin 1 and Map1lc3) expression in each group of cells.

**Results** Compared with the group I, target genes (Bclin 1 and Map1lc3) of the group III and the group IV had a significant decreased expression (p<0.01). The expression of the group IV were all lower than the group I (p>0.05). The group IV was better than the group III (p<0.05). Via TEM detection, the cells showed typical...
The study of derivation from bone marrow mesenchymal stem cells into cardiomyocyte-like cells in vitro via cardiotrophin-1

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Objective To investigate the effects of CT-1 on differentiation of induced swine BMSCs in vitro independently or with 5-aza.

Methods BMSCs were divided into four groups: bland control; induced with CT-1; induced with 5-aza; induced with 5-aza combined with CT-1. After 4 weeks of induced culturing, the differentiation of cardiomyocyte induced were estimated by cTnT and α-actinin.

Result Red fluorescence staining of α-actinin shows: the differentiation rate of myocardial cells of the composite group is greatest (36.500 ± 4.0927%); 5-aza group is greater than CT-1 group. Red fluorescence staining of cTnT shows: the differentiation rate of myocardial cells of the composite group is greatest (36.500 ± 4.0927%); 5-aza group is greater than CT-1 group; CT-1 group is greater than control.

Conclusions CT-1 can increasing the inducing rate combined with 5-aza.

The study of regulation of T-type Ca ion channel in lysophosphatidylcholine-stimulated cardiomyocytes

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Objectives To study the effect of lysophosphatidylcholine (LPC) in myocardial cells in T-type calcium channel currents (ICaT), and the hypothesis that LPC accumulation in intracellular and/or interstitial space in cardiomyocytes may underlie as a mechanism for tachyarrhythmias during cardiac ischaemia.

Methods Neonatal rat cardiomyocytes from 1 to 3-day-old Wistar rats and hypertrophied ventricular myocytes from Wistar rat were prepared. A single dose of 60 mg/Kg monocrotaline was injected into the intraperitoneal cavity at the age of 8 weeks old, and right nephrectomy was performed. A single dose of 60 mg/Kg monocrotaline was injected into the intraperitoneal cavity at the age of 8 weeks old, and right nephrectomy was performed.

Results (1) LPC markedly accelerated the spontaneous beating rates of neonatal rat cardiomyocytes from 42±8 bpm in control to 64±8 bpm after LPC application in 5 min at the physiological [Ca2+]i condition (pCa=7.2). (2) In neonatal cardiomyocytes, ICa,T was significantly increased by 10 μM LPC by 21.5% when [Ca2+]i was high (pCa=7). Intracellular Ca2+-dependent augmentation of ICa,T by LPC was confirmed not only in neonatal cardiomyocytes but in adult ventricular myocytes from the hypertrophied heart. In this experiment, ICa,T was significantly increased by 10 μM LPC by 23.5% when [Ca2+]i was high (pCa=7), although it was unchanged when [Ca2+]i was low (pCa=11). (3) LPC exerted no effect on the CaV3.1 T-type Ca2+ channel current (ICaV3.1) regardless of the [Ca2+]i condition at a pCa of 7 (solution F) or at a pCa of 11 (solution A). In contrast, LPC upregulated the CaV3.2 T-type Ca2+ channel current (ICaV3.2), which was much larger at a pCa of 7 than that at a pCa of 11. (4) A specific PKC inhibitor Ro-32-0452 completely blocked the effect of LPC on ICaV3.2. However, in the same culture condition, a specific PKCβ inhibitor G6 6976 (20 nM) and a specific PKCβII inhibitor C84353 (2 μM) did not modify the effect of LPC on ICaV3.2.

Conclusion Using atorvastatin, prior to the occurrence of induced autophagy, which may be related to the role of atorvastatin’s improvement on endothelial function. However, using atorvastatin, prior to the occurrence of induced autophagy, can not quite inhibit the occurrence of autophagy.