influx via NCX. Thus, YM-244769 has therapeutic potential as an efficient re-noprotective drug.

3367-Pos Board B228
Vascular Na⁺/Ca²⁺ Exchanger Type-I Contributes to Hypertension in Pseudohypaldosteronism Type II and Cushing’s Syndrome Models
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Pseudohyaldosteronism type II (PHAI) is an autosomal dominant disease characterized by hypertension due to increased renal salt reabsorption. Cush-
ing’s syndrome is associated with excessive cortisol secretion by ectopic ACTH-producing tumors and may result in hypertension. Here we examined the role of NCX (Ca¹⁺/Na⁺ exchanger type-I (NCX1)) in hypertension of these model mice using specific NCX inhibitors and genetically engineered mice. NCX in-
hibitors lowered arterial hypertension in IKd4 mutant knockin mice (present-
ing the phenotype of PHAI) and chronically ACTH-administered mice. Furthermore, homozygous knockout of NCX1 was resistant to development of hypertension in these model mice, whereas vascular overexpression of NCX1 accelerated their hypertension. Since NCX inhibitors reversed the cyto-
solic Ca²⁺ elevation and vasoconstriction induced by non-nanotmal ouabain, cir-
culating endogenous cardiac glycosides may be involved in hypertension of these model mice. Thus, vascular NCX1 contributes to hypertension in PHAI and Cush-
ing’s syndrome, and NCX inhibitors might be therapeutically useful for their hypertension.

3368-Pos Board B229
Reverse Mode of the Sodium Calcium Exchanger is Enhanced in Malig-
nant Hyperthermia Susceptible Skeletal Muscle
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Intracellular Ca²⁺ concentration [Ca²⁺i], and intracellular Na⁺ concentration [Na⁺i], were elevated in swine (RyR1-R163C) and rodent (RyR1-R163C) MH susceptible (MHS) compared to Wt (MHN) muscle fibers. In both MHS and MHS muscle fibers stepwise reduction of external [Na⁺i] gradually in-
ncreased [Ca²⁺i], which could be prevented by removal of extracellular Ca²⁺ ([Ca²⁺o]). Disruption of the T-system by glycerol treatment also reduced the magnitude of the [Ca²⁺i], elevation induced by Na⁺ free solution in both groups. Administration of KBR7943 reduced [Ca²⁺i], in MHS and MHS mus-
cle fibers, and ameliorated the magnitude of the elevation of [Ca²⁺i], observed during a MH episode. However, YM-244769 a NCX blocker that preferen-
tially inhibits NCX3 reverse mode did not reduce [Ca²⁺i], in either MHN or MHS muscle fibers at rest, but did reduce the amplitude of the elevation of [Ca²⁺i], induced by halothane in R163C MHS muscle fibers. These results sup-
port the existence of a functional NCX reverse mode in pig and mouse skeletal muscle, which appears to be enhanced in MHS muscle fibers, and ameliorated the magnitude of the elevation of [Ca²⁺i], in MHS muscle fibers during exposure to halothane.

3369-Pos Board B230
Atrial-Specific NCX KO Mice Reveal Dependence of Sinoatrial Node Pacemaker Activity on Sodium-Calcium Exchange
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The cardiac sodium-calcium exchanger (NCX1) is hypothesized to play a ma-
ajor role in sinoatrial node (SAN) pacemaker activity. To test this hypothesis, we used a novel technique to generate an atrial-specific knockout (KO) of NCX1 in mice using the sarcolipin promoter. At 12 weeks, there is no evidence of NCX1 in the atrium or SAN by immunostaining or immunoblot. KO mice exhibit atrial dilation and ventricular hypertrophy with mild ventric-
ular dysfunction. On electrocardiography, KO mice have no P waves and a rel-
tively slow junctional escape rhythm (231 ± 16 bpm, n = 4) compared to normal sinus rhythm in Wt (422 ± 63 bpm, n = 2, p = 0.01). Furthermore, recordings of cardiac electrograms in Langendorff-perfused hearts show no evidence of atrial activity in KO. In patch clamped SAN cells isolated from KO mice, there is no NCX activity in response to caffeine-induced SR Ca²⁺ release. L-type Ca²⁺ current is decreased in KO by ~50% but there is no significant difference in funny current (Iₜ) amplitude between WT and KO. Spontaneous action potentials (APs) are absent in KO, even after application of isoproterenol (ISO, 1µM). However, we were still able to evoke APs in patch clamped KO cells under current clamp conditions, indicating that KO cells are capable of electrically stimulated depolarization. The maximum di-
astolic potential (MDP) was slightly more depolarized in KO (-57 ± 2.0 mV) compared to WT (-70 ± 2.5 mV, p < 0.001), which could theoretically reduce spontaneous activity. However, reducing extracellular K⁺ to lower the MDP in KO to WT values failed to restore rhythmic pacemaker activity. Thus, we conclude that NCX1 is required for normal pacemaker activity in the murine SAN.

3370-Pos Board B231
A New View of Insulin Action
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Insulin not only stimulates the Na-K-pump but by stimulating the Na-H ex-
change pump increases intracellular pH, pHl. Fidelman et al (1982) found, as predicted by thermodynamic theory, the effect of insulin on glycolysis varied linearly with log[Na] and the insulin effect being converted from stimulation of glycolysis to a 51% inhibition at 0.12 nM Nao. Zierler & Rabinowitz (1964) demonstrated in the forearm of human males that levels of insulin too low to affect glucose uptake, as little as 38 micro units per cc was sufficient to stimulate the Na-K-pump. As reflected by increased po-
tassium uptake, while producing no effect upon glucose uptake. This one exper-
iment forces us to realize that the main action of insulin is not to regulate blood glucose levels, but to regulate the Na-K-pump and Na-H exchange. Since the Na-K-pump uses about 25% of the ATP production in a resting muscle, stimu-
lation of the Na-K-pump increases the consumption of ATP so much that even high levels of stimulation, it is necessary to get more glucose into the cell to manufacture more ATP. Since insulin decreases Na, elevates pH, and increases production of ATP, one would expect that the decreased insulin, such as seen in diabetes or in fasting, would result in the reverse of these changes. Lowering plasma levels of insulin in rats by small doses of streptozotocin or by fasting produced an increase in Nao of about 30%, a decrease in pH of 0.15 units, and a 24% decrease in ATP, indicating the insulin effect on the Na-K-pump is due to changes in the diuretics to cause type 2 diuretics has been shown to be due to their ability to cause potassium loss. Not surprisingly, diets with large amounts of potassium and small amounts of sodium have been shown to reverse diabetes.

3371-Pos Board B232
Platelet Activating Factor Stimulates Sodium-Proton Exchange
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Sodium-hydrogen exchanger (NHE), the principal sarcolemmal acid exchanger in ventricular myocytes is stimulated by a variety of autocrine/paracrine fac-
tors and contributes to myocardial injury and arrhythmias during ischemia/ reperfusion (IR). Platelet-activating factor (PAF, 1-O-alkyl-2-acetyl-sn-
glycero-3-phosphocholine) is a potent proinflammatory phospholipid that is released in the heart in response to oxidative stress and promotes myocardial IR injury. PAF stimulates NHE in neutrophils and platelets, but its effect on cardiac NHE (NHE1) is resolved. We utilized quiescent guinea pig ventricular myocytes bathed in bicarbonate-free solutions and used epifluorescence to mea-
sure intracellular pH (pHi). Methylcarbamyl-PAF (C-PAF, 200 nM), a meta-
bolically-stable analog of PAF, significantly increased steady-state pHi. The alkalosis induced by C-PAF was completely blocked by the NHE inhibitor, cariporide, and by sodium-free bathing solutions, indicating it was mediated by NHE activation. C-PAF also significantly increased the rate of acid xtrusion induced by intracellular acidosis. The ability of C-PAF to increase steady-state pHi was completely blocked by the PAF receptor inhibitor WEB 2086 (10 µM), indi-
cating the PAF receptor is required. A mitogen-activated protein (MAP) ki-
nase kinase (MEK) inhibitor (PD98059, 25 µM), completely blocked the rise in pHi induced by C-PAF, suggesting participation of the MAP kinase signaling cascade downstream of the PAF receptor. Inhibition of protein kinase C (PKC) with GF109203X (1 µM) and chelerythrine (2 µM) did not significan-
tly affect the alkalosis induced by C-PAF. In summary, these results provide evidence that: a) PAF stimulates cardiac NHE1, b) the effect occurs via the PAF receptor, and c) signal relay requires participation of the MAP kinase cascade.