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1	A Pharmacologic Activator of Endothelial KCa Channels Increases Systemic Conductance
2	and Reduces Arterial Pressure in an Anesthetized Pig Model
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30 Abstract

31 SKA-31, an activator of endothelial KCa2.3 and KCa3.1 channels, reduces systemic blood

- 32 pressure in mice and dogs, however, its effects in larger mammals are not well known. We
- therefore examined the hemodynamic effects of SKA-31, along with sodium nitroprusside
- 34 (SNP), in anesthetized, juvenile male domestic pigs. Experimentally, continuous measurements
- of left ventricular (LV), aortic and inferior vena cava (IVC) pressures, along with flows in the
- 36 ascending aorta, carotid artery, left anterior descending coronary artery and renal artery, were
- performed during acute administration of SKA-31 (0.1, 0.3, 1.0, 3.0 and 5.0 mg/ml/kg) and a
- single dose of SNP (5.0 μ g/ml/kg). SKA-31 dose-dependently reduced mean aortic pressure
- 39 (mP_{AO}), with the highest dose decreasing mP_{AO} to a similar extent as SNP (-23 \pm 3 and -28 \pm 4
- 40 mmHg, respectively). IVC pressure did not change. Systemic conductance and conductance in
- 41 coronary and carotid arteries increased in response to SKA-31 and SNP, but renal conductance
- 42 was unaffected. There was no change in either LV stroke volume (SV) or heart rate (*versus* the
- 43 preceding control) for any infusion. With no change in SV, drug-evoked decreases in LV stroke
- 44 work (SW) were attributed to reductions in mP_{AO} (SW vs. mP_{AO}, $r^2 = 0.82$, P < 0.001). In
- 45 summary, SKA-31 dose-dependently reduced mP_{AO} by increasing systemic and arterial
- 46 conductances. Primary reductions in mP_{AO} by SKA-31 largely account for associated decreases
- 47 in SW, implying that SKA-31 does not directly impair cardiac contractility.

- 49
- 50 Abbreviations: G, conductance; HR, heart rate; IVC, inferior vena cava; KCa channel, calcium-
- 51 activated K⁺ channel; ; mPAO, mean aortic pressure; mP_{IVC}, mean inferior vena caval pressure;
- 52 PBS, phosphate-buffered saline; P_{LVED}, left ventricular end-diastolic pressure; SKA-31,
- naphthol[1,2-d]thiazol-2-ylamine; SNP, sodium nitroprusside; SV, stroke volume; SVR,
- 54 systemic vascular resistance; SW, stroke work; Vol_D, volume of distribution

56 **1. Introduction**

The vascular endothelium plays a critical role in the regulation of blood pressure and 57 blood flow distribution by controlling the intraluminal diameter of conduit and small resistance 58 arteries. This dynamic regulation occurs via the activation of distinct vasodilatory mechanisms in 59 the endothelium that reduce contractile tone in the surrounding vascular smooth muscle, leading 60 to increased intraluminal diameter, arterial conductance and blood flow. Major pathways 61 contributing to endothelium-dependent vasodilation include the *de novo* synthesis of nitric oxide, 62 prostacyclin and the generation of a hyperpolarizing electrical signal that acts on vascular 63 smooth muscle. Endothelium-dependent hyperpolarization (EDH) is generated primarily via the 64 activation of endothelial small- and intermediate-conductance, Ca²⁺-activated K⁺ channels 65 (KCa2.3 and KCa3.1 channels, respectively) and is transmitted via myoendothelial gap junction 66 connections to the adjacent smooth muscle, where it causes membrane hyperpolarization and 67 reduced Ca²⁺ influx via voltage-gated Ca²⁺ channels. Small-molecule activators of KCa2.3 and 68 KCa3.1 channels evoke direct hyperpolarization of endothelial cells [1-5], relax myogenically 69 70 active resistance arteries [1,6] increase coronary flow in isolated heart preparations [7] and lower 71 blood pressure in normo- and hypertensive mice [2,5]. In conscious dogs, bolus administration of a KCa channel activator transiently lowers systemic blood pressure [4]. In contrast, genetic 72 73 knockout of endothelial KCa channels in mice leads to elevated systemic blood pressure and impairs or abolishes stimulus-evoked vasodilatory processes in isolated arteries and tissues [8]. 74 75 Endothelial KCa channel activity may also be important in disease settings, as a KCa channel activator is able to restore agonist-evoked vasodilatory responses in the coronary circulation of a 76 77 rodent model of type II diabetes exhibiting endothelial dysfunction [9].

To advance our knowledge of the in vivo cardiovascular effects of endothelial KCa 78 79 channel activators, the goal of the present study was to investigate the systemic hemodynamic effects of SKA-31, a recently described, second-generation KCa channel activator [2], in a large 80 animal model, the anesthetized, instrumented pig. Our results demonstrate that bolus intravenous 81 injections of SKA-31 dose-dependently lower mean aortic pressure and increase systemic 82 conductance to levels comparable to those elicited by the nitrovasodilator sodium nitroprusside 83 84 (SNP). SKA-31 increased arterial conductance in coronary and carotid arteries, indicating that SKA-31 may have broad vasodilatory action in the vasculature. Neither SKA-31 nor SNP 85 86 appeared to directly alter myocardial contractility. In summary, our data demonstrate that SKA-

peripheral circulation of the anesthetized pig. These observations suggest that SKA-31 may also

89 be an effective vasodilator in the human vasculature.

90

87

91 **2. Methods and Materials**

The experimental protocols used in this study were approved by the University of
Calgary Animal Care Committee, and conform to the NIH-published Guide for the Care and Use
of Laboratory Animals (8th edition, 2011), and are further consistent with those of the American
Physiological Society.

96

97 2.1 Animal preparation

Seven male domestic pigs (25-30 kg body weight, average weight 27 kg, 16-18 weeks of 98 age) were studied. Pigs were pre-medicated with an intramuscular injection of ketamine 99 hydrochloride (600 mg), fentanyl citrate (2 mg), and midazolam (10 mg). A 20-gauge catheter 100 was inserted into an ear vein and anesthesia was induced with sodium thiopental (25 mg/kg). 101 102 Anesthesia (level 3) was maintained with a continuous intravenous (I.V.) infusion containing a mixture of fentanyl citrate (0.04 mg/ml), midazolam (0.025 mg/ml) and ketamine hydrochloride 103 104 (0.3 mg/ml) at a rate of 100 ml/hour. Both isoflurane (less than 1% in the ventilator) and lidocaine (3 bolus intravenous administrations, 1 mg/kg, 5 min apart, followed by an I.V. 105 106 infusion of 0.75 - 1.0 mg/min) were used as required. The drug infusion rates were adjusted as 107 necessary to ensure deep sedation without spontaneous respiratory effort. The animals were 108 intubated with a cuffed endotracheal tube and ventilated with constant-volume ventilator (Harvard Apparatus, Millis, MA) with a 50% oxygen -50% nitrous oxide mixture. Tidal volume 109 110 and respiratory rate were adjusted to maintain physiological values of blood gases and pH in accordance with recommended ventilation parameters for large animals [10]. PaCO₂ was 111 112 maintained between 35 and 45 mmHg.

113 A median sternotomy was performed and the hearts were delivered from the pericardium 114 through a base-to-apex incision. Sonomicrometry crystals (Sonometrics, London, ON) were 115 implanted in the left ventricular endocardium and mid-wall of the septum to measure the minor-116 axis septum-to-left ventricular free wall and left ventricular antero-posterior dimensions [11-13]. 117 Ultrasonic flow probes (Transonic Systems, Ithaca, NY) were placed on the ascending aorta, 118 descending aorta (just above diaphragm), inferior vena cava (IVC) (just above the diaphragm), right carotid artery, left renal artery, and left anterior descending coronary artery. Thin walled 7-119 120 French fluid-filled catheters connected to pressure transducers (model P23 ID; Statham Gould, Oxnard, CA) were inserted into the left ventricle (LV) (P_{LV}; retrograde through the left carotid 121 artery), aorta (P_{AO}; retrograde through the right femoral artery) and IVC (P_{IVC}; through the right 122 jugular vein). An intravenous line was placed in the left external jugular vein for volume loading 123 124 (PentaspanTM, 10% pentastarch in 0.9% NaCl) to replenish fluid loss during surgery. A thinwalled catheter was connected to the intravenous line for bolus infusions. Arterial samples for 125 blood-gas analysis were obtained from a side-port on the aortic catheter. Body temperature was 126 monitored with a rectal thermometer. After instrumentation, the heart was returned to the 127 pericardium, which was closed with individual sutures, taking care not to compromise pericardial 128 volume [14]. A single-lead electrocardiogram (ECG) was recorded. 129

130

131 2.2 Experimental protocol

Simultaneous pressure, dimension and flow measurements were recorded at baseline and 132 during each intervention. After stabilization at an LV end-diastolic pressure (P_{LVED}) of ~10 133 mmHg (11 ± 1 mmHg), control data were collected for 60 s, immediately preceding a 5-min 134 135 recording period, during and after drug infusion. Each 20 ml infusion was delivered over a 60-s period and proceeded in ascending order of SKA-31 dosage (0.1, 0.3, 1.0, 3.0, and 5.0 mg/ml/kg) 136 137 followed by a single dosage of sodium nitroprusside (SNP; 5.0 µg/ml/kg). Washout and recovery periods of 15-20 min were interposed between drug infusions. At the end of the experiment, the 138 139 animals were sacrificed by a bolus KCl injection and the positions of the sonomicrometry crystals within the myocardium were verified. 140

141 SKA-31 was synthesized and tested for identity and purity (NMR and HPLC/MS) as previously described [2]. SKA-31 was dissolved in a vehicle solution comprised of Cremophor 142 EL (10% v/v) and phosphate-buffered saline (PBS) (90% v/v). Briefly, an aliquot of Cremophor 143 EL was first heated in a beaker on a magnetic stir plate to a temperature of $\sim 60^{\circ}$ C. The desired 144 amount of solid SKA-31 was then added to the heated Cremophor EL liquid as it was being 145 146 stirred. Once the added SKA-31 had dissolved completely, heating was stopped and stirring was maintained. The first few milliliters of PBS were then added slowly to the SKA-31/Cremophor 147 148 EL solution and the remaining amount was added more quickly. The final SKA-31 solution was

150 Solutions of SKA-31 in Cremophor-EL/PBS were freshly prepared for each experiment.

151

152 *2.3 Data analysis*

The conditioned signals were passed through a low-pass filter (100 Hz) and were 153 digitized and recorded at 100 Hz (Sonometrics Corp. acquisition system, London, ON). The 154 digitized data were analyzed on a personal computer using custom software (CV Works, 155 Calgary, AB) developed in our laboratory. Baseline and control data are expressed as mean 156 values for the 60-s period immediately preceding each infusion event. All data associated with 157 administration of drug or control solutions were extracted at the time of greatest decrease in 158 mP_{AO}. If mP_{AO} did not change by at least 5 mmHg during a given intervention, acquired data 159 points were averaged for the first 60 s of that period. 160

Systemic conductance (G_{systemic}, the reciprocal of systemic vascular resistance, SVR) was 161 calculated as mean aortic flow / $(mP_{AO} - mP_{IVC})$ and expressed as a percent change from the 162 preceding control value. Carotid conductance (G_{carotid}), renal conductance (G_{renal}) and coronary 163 164 conductance (G_{coronary}) were expressed similarly and calculated by respectively substituting mean carotid, renal and coronary flow for mean aortic flow. LV stroke work (SW) was calculated as 165 166 LV stroke volume (SV) x [mean P_{LV} (systolic) – P_{LVED}], where mean P_{LV} (systolic) was calculated as PAO (diastolic) + 2/3 [PAO (systolic) - PAO (diastolic)]. As an index of LV end-167 168 diastolic volume, LV area (A_{LVED}), was calculated as the product of the 2 minor-axis LV 169 dimensions [15,16]. SW and A_{LVED} values following drug infusions are expressed as the percent 170 change from the preceding control values determined using the same calculations.

171

172 2.4 Measurements of SKA-31 Concentration in Plasma

Blood samples (~2 ml) were taken via a catheter inserted into the left external jugular vein at various intervals following drug infusion at the same site. Samples were collected in heparinized tubes to prevent coagulation and centrifuged at 400 x g for 20 min at 4°C. The resulting supernatants were then stored at -80°C prior to analysis. Plasma samples were then processed and analyzed in duplicate by HPLC/MS as recently described [4] and SKA-31 concentrations were determined from a standard curve. A semi-logarithmic plot of SKA-31 plasma concentrations vs. time was fitted with the following equation:

180	SKA-31 Plasma Concent. = $C_0 * \exp(-k_e t)$
181	Where C_0 = the maximal initial concentration of SKA-31 in the plasma calculated from the y-
182	intercept of the fitted line, k_e is the rate constant and t is the time interval following SKA-31
183	infusion. The volume of SKA-31 distribution (Vol _D) was calculated as follows:
184	$Vol_D = SKA-31 \text{ dosage/}C_0$
185	
186	2.5 Statistical analysis
187	Statistical comparisons were performed using SigmaPlot (Systat Software, Inc. 2012). In
188	Figure 8, a linear correlation was calculated for the percentage changes for mP_{AO} and stroke
189	work during saline, vehicle, and all drug infusions ($y = y_0 + a * x$). The Student's paired <i>t</i> -test was
190	used to test for the significance of changes between a given infusion (i.e. vehicle or drug) and the
191	preceding control period. Repeated-measures ANOVA (Holm-Sidak method) was used to test for
192	the significance of differences between vehicle/SKA-31 infusions and SNP. A P value <0.05 was
193	considered statistically significant. Except where noted, data are presented as mean \pm SEM.

194

195 **3. Results**

Seven anesthetized, juvenile pigs were acutely implanted with blood pressure transducers and Doppler flow probes that allowed us to measure mean aortic and inferior vena cava pressures, systemic conductance and regional conductance in carotid, renal and coronary arteries. Myocardial performance was monitored via a single lead electrocardiogram and implanted sonomicrometry crystals in the myocardium to assess LV dimensions. Table 1 presents average hemodynamic parameters in all 7 animals measured at baseline, following instrumentation and recovery and before the first experimental infusion (saline).

203

204 *3.1 SKA-31 Dose Response*

Following surgical interventions, animals were allowed to recover until steady-state basal levels of mean aortic pressure and heart rate were achieved. After a minimum 10 min period of steady-state baseline recording, we commenced with the first (saline) infusion. Figure 1 displays representative tracings of the effect of individual bolus administrations of saline, drug vehicle, SKA-31 (0.1 - 5.0 mg/ml/kg) on mean aortic pressure (mP_{AO}, panel A), systemic conductance (panel B), measured conductance in carotid, coronary and renal arteries (panel C) and heart rate 211 (panel D). While SKA-31 infusions had clear effects on these hemodynamic parameters, neither 212 saline nor vehicle infusions had any observable effects. In an effort to benchmark the effects of SKA-31 on the measured parameters, we infused a single dose of the well characterized 213 nitrovasodilator sodium nitroprusside (SNP) following recovery from the SKA-31 evoked 214 hemodynamic changes. As displayed on the right hand side of Figures 1A-D, SNP infusion (5.0 215 $\mu g/ml/kg$) produced qualitatively similar changes in mP_{AO}, systemic conductance, carotid, 216 coronary and renal artery conductances and heart rate when compared with SKA-31. In case of 217 218 mP_{AO} (Fig. 1A), intravenous infusion of SKA-31 significantly decreased mP_{AO} in a dosedependent manner versus each preceding control period, with the greatest decrease occurring 219 after the highest dose (5.0 mg/ml/kg) (Figure 2). SNP infusion also significantly decreased mP_{AO} 220 and this change was comparable to that measured following infusion of 5.0 mg/ml/kg SKA-31. 221 222 As quantified in Figure 3, the time to peak response for the SKA-31 induced decrease in 223 mP_{AO} was slowest at 0.1 mg/ml/kg drug administration and became faster with increasing dosages. At a dosage of 5.0 mg/ml/kg, SKA-31 infusion resulted in a significantly faster decline 224

in mP_{AO} compared with SNP.

In contrast to the observed decreases in mP_{AO} , SKA-31 did not significantly alter mean inferior vena cava pressure (mP_{IVC}), compared with preceding control values (Figure 4). In the case of SNP, we did observe a trend toward lower mP_{IVC} , although this change did not reach statistical significance.

230

231 *3.2 Conductance and Resistance*

Figure 5 shows the effect of SKA-31 and SNP administration on absolute changes in 232 233 systemic vascular resistance (SVR) (Fig. 5A), along with the calculated percent changes in systemic conductance (Fig. 5B). We observed no changes in SVR versus the preceding control 234 values following infusions of saline, drug vehicle or lower dosages of SKA-31 (0.1 and 0.3 235 mg/ml/kg), whereas dosages of 1.0, 3.0 and 5.0 mg/ml/kg each significantly decreased systemic 236 237 resistance. SKA-31 at the highest dosage decreased SVR to a level comparable to that evoked by 5.0 µg/ml/kg SNP. Predictably, the inverse relationships were observed for drug-induced 238 changes in systemic conductance (Fig. 5B). 239

In addition to its impact on systemic conductance, we also examined the effect of SKA31 on blood flow in select vascular regions. As shown in Figures 1C and 6, SKA-31 and SNP

242 infusions produced qualitatively similar effects on conductance in the right carotid artery (G_{carotid}), left anterior descending coronary artery (G_{coronary}), and left renal artery (G_{renal}). SKA-31 243 increased G_{carotid} at dosages of 3.0 and 5.0 mg/ml/kg and produced a maximal change in 244 245 conductance similar to that observed with 5.0 μ g/ml/kg SNP. In the left anterior descending coronary artery, SKA-31 also significantly increased G_{coronary} at doses of 3.0 and 5.0 mg/ml; the 246 247 increase evoked by the latter dose approximated that observed with SNP. Interestingly, neither 248 SKA-31 nor SNP significantly increased blood flow in the renal artery (compared with the 249 preceding control conductance values) and vasodilatory responses in this artery were generally 250 blunted compared with carotid and coronary vessels (Fig. 1C).

251

252 *3.3 Cardiac Function*

253 As depicted in Figure 7A, infusions of SKA-31 and SNP did not produce significant 254 changes in either cardiac stroke volume (SV) or heart rate (HR) under our experimental conditions. We further examined potential drug-induced changes in the left ventricular end 255 diastolic area (A_{LVED}), as measured by sonomicrometry crystals implanted in the septal wall and 256 LV endocardium, and the calculated LV stroke work (SW_{LV}); both of these parameters are 257 expressed as the percent change from the respective preceding control value. Over the dosage 258 range of 1.0 to 5.0 mg/ml/kg, SKA-31 produced very modest decreases in ALVED (< 5% below 259 control), whereas SNP reduced A_{LVED} by an average of 8% compared with control. In contrast to 260 the slight decreases observed in ALVED, SKA-31 evoked a clear, dose-dependent reduction in 261 SW_{LV} over the range of 0.1 to 5.0 mg/ml/kg and produced a similar maximal decrease at a 262 dosage of 5.0 mg/ml/kg (-29%) as that observed following SNP infusion (-32%). 263

To evaluate in greater depth the observed decreases in SW_{LV} following SKA-31 and SNP 264 administrations, we plotted the calculated percent changes in SW_{LV} versus the observed percent 265 changes in mean aortic pressure (mP_{AO}). The scatter plot in Figure 8 shows the relation between 266 SW_{LV} and mP_{AO}, based on the pooled data derived from all infusions of saline, vehicle, SKA-31 267 and SNP for the 7 animals employed in our study. Importantly, the calculated r^2 value of 0.82 268 for the linear regression line indicates that more than 80% of the variance in SW_{LV} can be 269 270 explained by the variance in mP_{AQ} . Using a similar approach, we also plotted the percent changes in A_{LVED} versus mP_{AO} for all animals and infusions (i.e. saline, vehicle, drug) examined. 271 Linear regression analysis of this relation yielded a r^2 value of only 0.24, indicating that no more 272

than 25% of the variance in A_{LVED} can be explained by the variance in mP_{AO} (P<0.001; data not

shown). Based on these results and the fact that drug infusions did not change stroke volume

under any condition (see Figure 7A), we conclude that neither SKA-31 nor SNP directly

276 impaired cardiac contractility.

277

278 3.4 Plasma Concentrations of SKA-31 Following Acute Infusion

In a separate group of 3 anesthetized and instrumented animals, we analyzed the plasma 279 concentrations of SKA-31 at select time points following acute intravenous infusion of a 3.0 280 mg/ml/kg SKA-31 bolus dose. Blood samples were withdrawn from the left jugular vein at ~1 281 min, 35 min and 75 min following complete infusion of the drug. The average free plasma 282 concentration of SKA-31 measured at each of the above time points (n = 3) was 77.5 \pm 27.7 μ M, 283 $27.3 \pm 3.8 \mu$ M and $27.4 \pm 4.9 \mu$ M, respectively. The volume of distribution for SKA-31 284 285 calculated from a semi-logarithmic plot of average SKA-31 plasma concentrations versus sampling times was 0.19 L/kg. 286

287

288 **4. Discussion**

289 Using an anesthetized and instrumented porcine model, we have provided the first 290 detailed description of the systemic hemodynamic actions of SKA-31, a small molecule activator of KCa 2.x and 3.1 channels [2], on key cardiovascular parameters in a large animal and how 291 292 these actions compare with those of SNP, an established nitrovasodilator and blood pressurelowering agent. As shown in Figures 1 and 2, intravenous administration of SKA-31 dose-293 294 dependently evoked significant decreases in mean aortic pressure, with the highest dose utilized in our study (5.0 mg/ml/kg SKA-31) producing a similar decrease in mPAO as that observed with 295 SNP (-23±3 and -28±4 mmHg, respectively) (Fig. 2B). In previous studies [2,5,8], acute in vivo 296 297 administration of SKA-31 was shown to lower blood pressure in both normotensive and hypertensive mice and, more recently, Köhler and colleagues [4] have reported that acute 298 infusion of SKA-31 (0.4 and 2.0 mg/kg) transiently decreases systemic blood pressure in 299 conscious dogs. We also noted that the decrease in mPAO evoked by 5.0 mg/ml/kg SKA-31 was 300 301 more rapid compared with SNP (Fig. 3), even though both agents lowered mean aortic pressure to a similar extent (Fig. 2). The slower time course of the SNP-mediated drop in mP_{AO} may 302 303 reflect the fact that SNP requires vascular conversion/decomposition to release nitric oxide and

induce subsequent cellular actions in arterial smooth muscle [17], while SKA-31 directly
hyperpolarizes the endothelium by activating KCa channels. Collectively, these observations are
in agreement with the reported vasodilatory actions of SKA-31 in the intact coronary [7] and
skeletal muscle circulations [5,8] of rodents and the systemic circulation of the dog [4].

SKA-31 had no significant effect on mean inferior vena cava pressure (mP_{IVC}) (Fig. 4). In 308 the case of SNP, we did observe a trend towards lower mP_{IVC}, which would be in agreement with 309 310 the known clinical effects of SNP to lower central venous pressure, due to its ability to increase venous capacitance [18]. One possible reason for our observation is that mP_{IVC} was already quite 311 low under basal experimental conditions (~8 mmHg) and a further drug-induced decrease in 312 mP_{IVC} may have been difficult to detect in our anesthetized pigs. Although KCa2.3 and KCa3.1 313 channel mRNA and whole cell K⁺ currents have been reported in venous endothelial cells (e.g. 314 HUVECs) [1,3,19], we are unaware of data describing a direct vasodilatory effect of KCa 315 channel activators on veins or the venous circulation. 316

317 SKA-31 dosages of 1.0 to 5.0 mg/ml/kg increased systemic arterial conductance, with the highest dose producing an increase in conductance similar to that induced by SNP (Figure 5B). 318 319 We also observed increases in both carotid and coronary arterial conductances at 3.0 and 5.0 mg/ml/kg SKA-31 (Figure 6), which were similar to those observed with SNP at the highest 320 321 dosage of SKA-31. Interestingly, renal conductance appeared to be unaffected by either SKA-31 or SNP. In the case of SNP, this is somewhat unexpected, as other investigators have reported 322 323 that renal arteries are sensitive to nitrovasodilators [20,21]. The renal microcirculation is known to exhibit strong autoregulatory behavior [22,23], which is critical for ensuring adequate blood 324 325 flow to glomerular units and protecting them from arterial pressure-induced damage. One possible explanation for this apparent insensitivity of the renal conductance to SKA-31 and SNP 326 327 is that the renal circulation may have already been near-maximally dilated, due to a combination of intrinsic autoregulation and the somewhat lower mP_{AO} present in our anesthetized pigs. 328 Alternatively, it is possible that reduced arterial resistance triggered an increase in peripheral 329 330 sympathetic tone to counteract reduced blood pressure, which then limited renal arterial dilation. However, this possibility is less likely, as we observed no concomitant increase in heart rate with 331 332 declines in mP_{AO}, which one would anticipate with the activation of a baroreceptor feedback mechanism acting on the heart. 333

As small-conductance Ca²⁺-activated K⁺ channels have been reported in the atria and 336 337 pacemaker/conducting cells of murine and human cardiac tissue [24-27] and thus may be activated in response to systemic SKA-31 administration, we recorded various indices of 338 myocardial performance during SKA-31 infusions. Importantly, we observed no significant 339 change in left ventricular stroke volume following administration of either SKA-31 or SNP. In 340 contrast to Köhler and coworkers [4], who reported a pronounced increase in heart rate (HR) 341 following acute SKA-31 infusion, we did not detect a significant change in HR in response to 342 SKA-31 or SNP infusions in the anesthetized pig (Fig. 7). The difference in HR responses in 343 these two studies could be attributed to the difference in experimental models, as Köhler and 344 colleagues examined conscious dogs (presumably with unsuppressed baroreceptor reflexes 345 346 providing autonomic nerve input to the heart) versus our anesthetized, instrumented pig model. Importantly, the absence of SKA-31 induced changes in HR observed in our study strongly 347 suggests that plasma levels of SKA-31 sufficient to evoke substantial decreases in blood pressure 348 do not directly impact either pacemaker function or action potential propagation in the heart, as 349 350 revealed under conditions of minimal baroreceptor reflex activity.

Neither SKA-31 nor SNP significantly reduced central venous pressure (Fig. 4). One 351 352 possible explanation is that the relative magnitude of the arterial and venous effects of these vasodilatory agents may differ [18,28-30] or an increase in total venous capacitance may have 353 354 been limited by a modest elevation in arterial capacitance as a result of evoked vasodilation. Hemodynamically, a minor, undetected rise in total venous capacitance could explain the slight 355 356 decrease we observed in A_{LVED}, our measure of left ventricular end-diastolic volume, in response to SKA-31 and more so to SNP (Fig. 7B). Since left ventricular stroke work (SW) is a function 357 358 of both ventricular volume and pressure, the decreases observed in SW following administration of SKA-31 and SNP could be explained, in part, by the minor reduction in ALVED. However, 359 360 further analysis of these data clearly showed that changes in ALVED accounted for less than 25% of the variance in SW, whereas changes in mP_{AO} accounted for more than 80%. Thus, the 361 observed decreases in SW could be largely attributed to the reductions in mPAO associated with 362 363 drug administration (Figure 8). Furthermore, the observed reductions in mP_{AO}, indicative of left ventricular afterload, would be expected to offset the slight decreases in ALVED observed with 364

SKA-31 and SNP administrations, and the balance of these two effects would tend to maintain
stroke volume near control levels in the presence of either SKA-31 or SNP (Fig. 7A).

367 Our attempt to explore the pharmacokinetic behavior of SKA-31 revealed that its plasma levels measured 35-75 min following intravenous infusion of a 3.0 mg/ml/kg dose were higher 368 than the reported EC₅₀ values of SKA-31 for KCa3.1 channels (~0.3 µM) and KCa2.3 channels 369 $(\sim 2 \mu M)$ [2], suggesting that sustained activation of these endothelial channels might be 370 anticipated. However, the absence of prolonged hypotension following administration of SKA-371 31 in our anesthetized pigs suggests that the relationship between the free plasma concentration 372 of SKA-31 (plasma protein binding of SKA-31 in mice and dogs is reported to be 35-40%) [2,4] 373 and its vasodilatory actions may not be a direct one and may be complicated by the availability 374 375 of additional drug binding sites or a more complex whole body distribution pattern.

376 Another explanation for the relatively short-lived hemodynamic response following 377 SKA-31 infusion could be a "desensitization" of the pharmacological targets for SKA-31 actions. Both KCa3.1 and KCa2.3 channels are subject to regulation by intracellular second 378 379 messengers or protein kinases and phosphatases. For example, phosphorylation of channelassociated calmodulin by casein kinase 2 reduces the affinity of KCa2 channels for the 380 381 membrane phospholipid PIP₂ [31] and likely contributes to the inhibition of KCa2 channel 382 activity by Gq-associated G-protein coupled receptors. KCa3.1 activity is increased by phosphorylation of His358 in the channel's C-terminus through the histidine kinase nucleoside 383 diphosphate kinase B (NDPK-B) [32], while the PI₃P phosphatase myotubalarin related protein 6 384 (MTMR6) and the histidine phosphatase phosphohistidine phosphatase-1 (PHPT-1) inhibit 385 KCa3.1 function in T-cells [33,34]. Additionally, KCa3.1 currents can be regulated by cAMP-386 dependent protein kinase (PKA) via Ser phosphorylation sites in the C-terminus [35-37], which 387 may impair endothelium-dependent vasodilation [38]. 388

Finally, the ability of a KCa channel activator to lower blood pressure more effectively in hypertensive *versus* normotensive mice [2,5] suggests that this class of compound may be beneficial in the acute or chronic treatment of elevated blood pressure. Our results showing that SKA-31 reduces blood pressure and increases systemic conductance in the pig suggest that translational studies examining the potential blood pressure-lowering actions of a KCa channel activator in a large animal model of hypertension or vascular disease are likely feasible.

396 *4.2 Summary and Conclusions*

397 The results of our study demonstrate that the KCa channel activator SKA-31 effectively 398 and reversibly increases systemic conductance in a dose-dependent manner and lowers mean aortic pressure in a large animal model. The observed hemodynamic actions of SKA-31 closely 399 400 mimic those evoked by SNP. SKA-31 did not directly affect cardiac contractility, nor did it appear to impact heart rate or excitability. The common and overlapping cardiovascular 401 responses to SKA-31 and SNP are consistent with the conclusion that SKA-31 acts primarily on 402 blood vessels to evoke its effects on the systemic vasculature. Given that SKA-31 is strictly an 403 endothelium-dependent vasodilator [6], endothelial KCa channel activators may be useful as an 404 alternative pharmacologic strategy to evoke acute arterial vasodilation in situations where the 405 hemodynamic actions of SNP may not be desirable or effective (e.g. nitrate tolerance). 406 407 Acknowledgements 408 409 The authors would like to acknowledge the excellent surgical expertise of Ms. Cheryl Meek throughout this study. This work was supported by research funding to A.P. Braun (Canadian 410 411 Institutes of Health Research MOP 97901), to H. Wulff (National Institutes of Health NS072585) and to J.V. Tyberg (Kidney Foundation of Canada/Pfizer Canada). 412 413 Conflict of Interest: On behalf of the all authors, the corresponding author states that no 414 415 conflicts of interest exist. 416 417 References 418 [1] Sheng J-Z, Ella S, Davis MJ, Hill MA, Braun AP. Openers of SK_{Ca} and IK_{Ca} channels 419 enhance agonist-evoked endothelial nitric oxide synthesis and arteriolar dilation. FASEB 420 J 2009;23: 1138-1145. doi: 10.1096/fj.08-120451 421 422 [2] Sankaranarayanan A, Raman G, Busch C, Schultz T, Zimin PI, Hoyer J et al. Naphthol[1,2-d]thiazol-2-ylamine (SKA-31), a new activator of KCa2 and KCa3.1 423 potassium channels, potentiates the endothelium-derived hyperpolarizing factor response 424 and lowers blood pressure. Mol Pharmacol 2009;75: 281-295. doi: 425

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538

Table 1 – Baseline hemodynamic parameters in anesthetized pigs immediately prior to the
control saline infusion at the start of the experiment.

541

HR (bpm)	122±9
SV (ml)	23±3
P _{LVED} (mmHg)	11±1
mP _{AO} (mmHg)	71±6
mP _{IVC} (mmHg)	7±1

542

543 HR, heart rate in beats per minute; SV, left ventricular stroke volume; P_{LVED} , end-diastolic left 544 ventricular pressure; m P_{AO} , mean aortic pressure; m P_{IVC} , mean inferior vena cava pressure. Data 545 represent the means ± SEM calculated from 7 pigs in total.

- 546
- 547

548 **Figure Legends**

Figure 1 – Representative data from one pig demonstrating the rapid and reversible effects of 549 SKA-31 and sodium nitroprusside (SNP) following acute intravenous infusion on mean aortic 550 pressure (mP_{AO}) (panel A), systemic vascular conductance (panel B), measured conductance in 551 coronary, carotid and renal arteries (panel C) and heart rate (panel D). In each panel, the sections 552 of continuous data points displayed represent 5-min epochs that were extracted from the master 553 data record and illustrate the basal levels and evoked changes in the measured parameters in 554 555 response to the infusions. The horizontal bars and labels provided beneath each panel specify the 556 experimental infusion for the 5-min section of data appearing immediately above each 557 description. Note that all displayed data were acquired simultaneously during the experiment. 558 Individual infusions were separated by a 15-20 min recovery period (indicated by the breaks 559 between the sections of data points) and control hemodynamic data were acquired for the first 1-2 min period immediately prior to a given infusion, once a steady baseline was clearly apparent 560 561 (not shown).

- Figure 2 Quantification of mean aortic pressure (mP_{AO}) under control conditions and following acute infusion of SKA-31 (0.1 - 5.0 mg/ml/kg) and SNP ($5.0 \mu \text{g/ml/kg}$) (panel A). Panel B quantifies the drug-evoked changes in mP_{AO} relative to the preceding control value for each experimental condition. N = 7 animals for both panels A and B.
- 568 Figure 3 Quantification of the time to maximal change in mean aortic pressure (mP_{AO})
- following intravenous infusion of either SKA-31 (0.1 5.0 mg/ml/kg) or SNP (5.0 μg/ml/kg).
- 570 Administration of either saline or drug vehicle did not evoke measurable changes in mP_{AO} . The
- response evoked by 5.0 mg/ml/kg SKA-31 was significantly faster than that elicited by SNP, as
- determined by two-way ANOVA; P < 0.05, n = 7 animals.
- 573

Figure 4 – Lack of effect of SKA-31 (0.1 - 5.0 mg/ml/kg) on mean inferior vena cava pressure (mP_{IVC}) following acute administration. Histogram displays mP_{IVC} values recorded in response to infusions of saline, drug vehicle and the indicated dosages of SKA-31 and SNP. Values for baseline mP_{IVC} (control) immediately preceding each infusion are designated by the black bars.

Figure 5 – Acute administration of SKA-31 and sodium nitroprusside (SNP) reduce systemic
vascular resistance (SVR). Panel A displays absolute values for SVR recorded prior to a given
drug infusion and following SKA-31 and SNP infusions at the indicated dosages. For the latter
data, measurements were taken during the peak change in SVR. Panel B displays the calculated
percentage change in systemic vascular resistance under each infusion condition compared with
the preceding control.

585

Figure 6 – Quantification of evoked changes in arterial conductance calculated for the carotid, coronary and renal arteries in response to infusions of saline, drug vehicle, SKA-31 (0.1 - 5.0mg/ml/kg) and SNP (5.0μ g/ml/kg). Histogram displays the percentage change in conductance in each artery evoked by administered drugs relative to the preceding control value for each infusion. Asterisks indicate a statistically significant difference compared with the baseline conductance value preceding a given infusion.

Figure 7 – Quantification of the effects of acute administration of saline, drug vehicle, SKA-31 (0.1 - 5.0 mg/ml/kg) or SNP ($5.0 \mu \text{g/ml/kg}$) on left ventricular stroke volume and heart rate (panel A). No significant changes were noted for either stroke volume or heart rate in response to a given infusion compared with the values measured during the preceding control period. The histogram in panel B displays the percentage changes in left ventricular area (A_{LVED}) and stroke work (SW), relative to the baseline values measured during the control period preceding each indicated infusion.

600

601 Figure 8 – Scatter plot displaying the relation between observed changes in left ventricular stroke

work (SW) and mean aortic pressure (mP_{AO}) following infusions of saline, vehicle, SKA-31 and

603 SNP. Percent changes in mP_{AO}, along with accompanying percent changes in SW, were first

calculated in response to each infusion utilized in a given experiment. Data points from all 7

animals were then plotted against each other in a pair-wise fashion, as depicted by the individual

symbols on the graph. The straight line through the symbols represents a linear regression fit to

607 the pooled data points (r^2 value = 0.82; P < 0.001).

Table 1- Hemodynamic parameters at baseline, immediately prior to the saline infusion.

HR (bpm)	122±9
SV (ml)	23±3
P _{LVED} (mmHg)	11±1
mP _{AO} (mmHg)	71±6
mP _{IVC} (mmHg)	7±1

HR, heart rate in beats per minute; SV, left ventricular stroke volume; P_{LVED} , end-diastolic left ventricular pressure; mP_{AO}, mean aortic pressure; mP_{IVC}, mean inferior vena cava pressure. Data represent the means ± SEM calculated from 7 pigs in total.



Figure 2 Click here to download high resolution image







Figure 5 Click here to download high resolution image



В



Figure 7 Click here to download high resolution image





