A computational model predicts sex-specific responses to calcium channel blockers in mammalian mesenteric vascular smooth muscle Gonzalo Hernandez-Hernandez, Samantha C. O'Dwyer, Collin Matsumoto, Mindy Tieu, Zhihui Fong, Pei-Chi Yang, Timothy J. Lewis[&], L. Fernando Santana*, and Colleen E. Clancy*^ Department of Physiology & Membrane Biology, ^Center for Precision Medicine and Data Science, University of California School of Medicine, Davis, California, 95616 [&] Department of Mathematics, University of California, Davis, California, 95616 *Correspondence to: ceclancy@ucdavis.edu and lfsantana@ucdavis.edu Running title: sex-specific models of arterial myocytes

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47 Abstract

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49 The function of the smooth muscle cells lining the walls of mammalian systemic arteries 50 and arterioles is to regulate the diameter of the vessels to control blood flow and blood 51 pressure. Here, we describe an *in-silico* model, which we call the "Hernandez-Hernandez 52 model", of electrical and Ca²⁺ signaling in arterial myocytes based on new experimental 53 data indicating sex-specific differences in male and female arterial myocytes from murine 54 resistance arteries. The model suggests the fundamental ionic mechanisms underlying membrane potential and intracellular Ca²⁺ signaling during the development of myogenic 55 56 tone in arterial blood vessels. Although experimental data suggest that Kv1.5 channel 57 currents have similar amplitudes, kinetics, and voltage dependencies in male and female 58 myocytes, simulations suggest that the $K_V 1.5$ current is the dominant current regulating 59 membrane potential in male myocytes. In female cells, which have larger Kv2.1 channel 60 expression and longer time constants for activation than male myocytes, predictions from 61 simulated female myocytes suggest that $K_{V}2.1$ plays a primary role in the control of membrane potential. Over the physiological range of membrane potentials, the gating of 62 a small number of voltage-gated K⁺ channels and L-type Ca²⁺ channels are predicted to 63 drive sex-specific differences in intracellular Ca²⁺ and excitability. We also show that in 64 an idealized computational model of a vessel, female arterial smooth muscle exhibits 65 heightened sensitivity to commonly used Ca²⁺ channel blockers compared to male. In 66 summary, we present a new model framework to investigate the potential sex-specific 67 68 impact of anti-hypertensive drugs.

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70 Introduction

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72 Our primary objective was to develop and implement a novel computational model that 73 comprehensively describes the essential mechanisms underlying electrical activity and 74 Ca²⁺ dynamics in arterial myocytes. We aimed to uncover the key components necessary 75 and sufficient to fully understand the behavior of arterial vascular smooth muscle 76 myocytes and the cellular response to variations in pressure. The model represents the 77 first-ever integration of sex-specific variations in voltage-gated Kv2.1 and Cav1.2 78 channels, enabling the prediction of sex-specific disparities in membrane potential and 79 the regulation of Ca²⁺ signaling in smooth muscle cells from systemic arteries. To further 80 investigate sex-specific responses to antihypertensive medications, we extended our 81 investigation to include a one-dimensional (1D) representation of tissue. This approach enabled us to simulate and forecast the effects of Ca²⁺ channel blockers within the 82 controlled environment of an idealized mesenteric vessel. It is worth noting that this 83 84 computational framework can be expanded to predict the consequences of antihypertensive drugs and other perturbations, transitioning seamlessly from single-cell 85 86 to tissue-level simulations.

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Previous mathematical models^{1–4} of vascular smooth muscle myocytes generated to 88 describe the membrane potential and Ca²⁺ signaling in vascular smooth muscle cells 89 90 have described the activation of G-protein-coupled receptors (GPCRs) by endogenous or 91 pharmacological vasoactive agents activating inositol 1,4,5-trisphosphate (IP₃) and 92 ryanodine (RyR) receptors resulting in the initiation of calcium waves. Earlier models have also provided insights into the contraction activation by agonists and the behavior 93 94 of vasomotion. In a major step forward, the Karlin model⁵ incorporated new cell structure data and electrophysiology experimental data in a computational model that predicted the 95 essential behavior of membrane potential and Ca²⁺ signaling arising from intracellular 96 97 domains found in arterial myocytes. One notable limitation of earlier models is that they 98 are based entirely on data from male animals. Furthermore, many data used to 99 parameterize the Karlin model were obtained from smooth muscle from cerebral arteries. 100 While cerebral arteries are important for brain blood flow, they do not control systemic

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blood pressure. Furthermore, they do not take into consideration the role of $K_{V2.1}$ channels in the regulation of smooth muscle cell membrane potential.

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104 The function of the smooth muscle cells that wrap around small arteries is to regulate the 105 diameter of these vessels. Arterial myocytes contract in response to increases in 106 intravascular pressure⁶. Based on work largely done using cerebral arterial smooth 107 muscle, a model has been proposed in which this myogenic response is initiated when membrane stretch activates Na⁺-permeable canonical TRPC6^{7,8} and melastatin-type 108 109 TRPM4^{9,10}. A recent study in smooth muscle from mesenteric arteries identified two 110 additional TRP channels to the chain of events that link increases in intravascular 111 pressure to arterial myocyte depolarization: TRPP1 (PKD1) and TRPP2(PKD2) channels^{11,12}. Together, these studies point to an elaborate multiprotein complex that 112 plays a critical role in sensing pressure and initiating the myogenic response by inducing 113 114 membrane depolarization and activating voltage-gated, dihydropyridine-sensitive L-type Cav1.2 Ca²⁺ channels^{13,14}. Ca²⁺ entry via a single or small cluster of Cav1.2 channels 115 produces a local increase in intracellular free Ca²⁺ ([Ca²⁺]_i) called a "Ca_V1.2 sparklet"¹⁵⁻ 116 ¹⁸. Activation of multiple Ca_V1.2 sparklets produces a global increase in $[Ca^{2+}]_i$ that 117 118 activates myosin light chain kinase, which initiates actin-myosin cross-bridge cycling and thus contraction¹⁹. 119

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Negative feedback regulation of membrane depolarization and Ca²⁺ sparklet activation 121 occurs through the activation of large-conductance, Ca²⁺-activated K⁺ (BK_{Ca}) channels as 122 well as voltage-dependent Ky2.1 and Ky1.5/1.2 K⁺ channels²⁰⁻²³. BK_{Ca} channels are 123 124 organized into clusters along the sarcolemma of arterial myocytes²⁴ and are activated by Ca²⁺ sparks resulting from the simultaneous opening of ryanodine receptors type 2 125 126 (RyR2) located in a specialized junctional sarcoplasmic reticulum (SR) ^{22,25–28}. Because the input resistance of arterial myocytes is high^{29,30} (about 2-10 G Ω), even relatively small 127 128 currents (10-30 pA) produced by the activation of a small cluster^{22,31,32} of 6-12 BKca channels by a Ca²⁺ spark can transiently hyperpolarize the membrane potential of these 129 130 cells by 10-30 mV. Accordingly, decreases in BK_{Ca}, K_V1.2, K_V1.5, and/or K_V2.1 channels

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131 depolarize arterial myocytes, increasing Ca $_{v}$ 1.2 channel activity, [Ca²⁺]_i, and contraction 132 of arterial smooth muscle^{21,33–36}.

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A recent study by O'Dwyer et al.²⁰ suggested that K_V2.1 channels have dual conducting 134 135 and structural roles in mesenteric artery smooth muscle with opposing functional 136 consequences. Conductive Kv2.1 channels oppose vasoconstriction by inducing 137 membrane hyperpolarization. Paradoxically, by promoting the structural clustering of the Cav1.2 channel, Kv2.1 enhances Ca²⁺ influx and induces vasoconstriction. Interestingly, 138 139 $K_{V2.1}$ protein is expressed to a larger extent in female than in male arterial smooth 140 muscle. This induced larger Cav1.2 clusters and activity in female than in male arterial 141 myocytes.

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Here, we describe a new model, which we call the "Hernandez-Hernandez model", of 143 mesenteric smooth muscle myocytes that incorporates new electrophysiological and Ca2+ 144 145 signaling data suggesting key sex-specific differences in male and female arterial 146 myocytes. The model simulates membrane currents and their impact on membrane potential as well as local and global [Ca²⁺]; signaling in male and female myocytes. The 147 148 Hernandez-Hernandez model predicts that $K_V 2.1$ channels play a critical, unexpectedly 149 large role in the control of membrane potential in female myocytes compared to male 150 myocytes. Importantly, our model predicts that clinically used antihypertensive Cav1.2 151 channel blockers cause larger reductions in Cav1.2 currents in female than in male 152 arterial myocytes.

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Finally, we present a one-dimensional (1D) vessel representation of electrotonically coupled arterial myocytes connected in series. Predictions from the idealized vessel suggest that Ca^{2+} channel blockers are more potent in females resulting in a more substantial $[Ca^{2+}]_i$ reduction in female arterial smooth muscle compared to male. The Hernandez-Hernandez model demonstrates the importance of sex-specific differences in Cav1.2 and Kv2.1 channels and suggests the fundamental electrophysiological and Ca^{2+} linked mechanisms of the myogenic tone. The model also points to testable hypotheses

- 6
- 161 underlying differential sex-based pathogenesis of hypertension and distinct responses to
- 162 antihypertensive agents.
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166 **RESULTS**

In this study, we developed a computational model of the electrical activity of an isolated vascular smooth muscle cell (**Figure 1**). A key goal was to optimize and validate the model with experimental data and then use the model to predict the effects of measured sex-dependent differences in the electrophysiology of smooth muscle myocytes.

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172 In constructing the model, we first set out to measure the kinetics of the voltage-gated Ltype Cav1.2 currents (I_{Ca}) in male and female myocytes using Ca²⁺ as the charge carrier 173 174 as shown in Figure 2. These data provided information on the kinetics of Ca²⁺-dependent 175 activation and inactivation of Ica. Ica is critical in determining cytosolic concentration [Ca²⁺]_i 176 in vascular mesenteric smooth muscle cells and is the predominant pathway for Ca²⁺ entrv^{13,15,16,18,28,52}. Experiments using whole-cell patch-clamp were undertaken to 177 178 measure the time constants of activation and deactivation (panel 2A) and inactivation 179 (panel 2B) in male and female mesenteric artery smooth muscle cells shown as black 180 and blue symbols, respectively. While the data from male (n = 10) and female (n = 12)181 myocytes showed comparable activation time constants, there was an observable trend 182 of faster inactivation in the female cells in the lower voltage range, but the differences 183 were not statistically significant. Steady-state activation and inactivation were also 184 measured as shown in **panel 2C**, with male data in black symbols and female as blue 185 symbols. No observable differences in the gating characteristics of the male and female 186 I_{Ca} were measured. Finally, the current-voltage relationship is shown from measurements in female (blue) and male (black) in **panel 2D.** This analysis suggests that the amplitude 187 188 of I_{Ca} was larger in female than in male myocytes over a wide range of membrane 189 potentials.

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We next used the experimental measurements to build and optimize a Hodgkin-Huxley model based on the data described above. The model includes voltage-dependent activation and inactivation gating variables, dL and dF, respectively. We modeled both gates following the approach by Kernik *et al.* ⁴⁵. It is important to note that smooth muscle cells operate within a voltage regime defined by the window current, which ranges between -45 mV and -20 mV. Under these conditions, $[Ca^{2+}]_i$ remains below 1 μ M.

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197 Therefore, we did not consider the Ca²⁺-dependent inactivation gating mode of the channel^{2,53}.

199

200 The model of I_{Ca} is described by:

201

202 (1)
$$I_{Ca} = P_{Ca} * dL * dF * \frac{z_{Ca}^2 F^2 V}{RT} \left(\frac{[Ca]_i e^{\frac{Z_{Ca} F V}{RT}} - [Ca]_{out}}{e^{\frac{Z_{Ca} F V}{RT}} - 1} \right)$$

203

Where P_{Ca} is the ion permeability, R is the gas constant, F is the Faraday's constant, and z_{Ca} is the valence of the Ca²⁺ ion. Parameters were optimized to male and female experimental data as shown for activation time constants (T_{activation}) and inactivation time constants (T_{inactivation}) as solid lines in **Figure 2A** and **Figure 2B**, respectively. Model optimization to male and female activation and inactivation curves are shown in **Figure 2C**. The model was also optimized to the I_{Ca} current-voltage (I-V) relationships shown as solid lines in **Figure 2D**.

211

212 We next set out to determine sex differences in voltage-gated K⁺ currents (I_{K}) in male and 213 female mesenteric smooth muscle cells. Ik is produced by the combined activation of Kv and BK_{Ca} channels. Following the approach previously published by O'Dwyer et al.,²⁰ we 214 quantified the contribution of K_V (I_{KV}) and BK_{Ca} (I_{BKCa}) current to I_K . K⁺ currents were 215 216 recorded before and after the application of the channel blocker iberiotoxin (IBTX; 217 100nm). Once identified the contribution I_{BKCa} current, we isolated the voltage-gated 218 potassium currents (I_{KV}) whose contributors include the voltage-gated potassium 219 channels Kv1.5 and Kv2.1. The presumed function of Kv1.5 and Kv2.1 channels on 220 membrane potential is to produce delayed rectifier currents to counterbalance the effect of the inward currents^{19,20}. 221

222

Having isolated I_{KV} , $K_V2.1$ currents were identified using the application of the $K_V2.1$ blocker ScTx1 (100 nM). As a result, the remaining ScTx1-insensitive component of the I_{KV} current was attributed to $K_V1.5$ channels. The results are shown in **Figure 3**. Experiments using whole-cell patch-clamp were undertaken to measure the steady-state

227 activation G/G_{max} of the K_V2.1current ($I_{KV2.1}$) as shown in **panel 3A** in female (blue) and 228 male (black) myocytes and no significant differences were observed. Measurements of 229 time constants of activation (**panel 3B**) of $I_{Kv2.1}$ in the voltage range of -30 to +40 mV in 230 female (blue, n=10) and male (black, n=7) myocytes exhibited significant differences. 231 Notably, activation time constants in male myocytes were smaller than those in female 232 myocytes, corresponding to a faster activation rate in males. The current-voltage 233 relationship of $I_{Kv2.1}$ is shown from measurements in female (blue, n=20) and male (black, 234 n=10) myocytes in **panel 3C**. Significant differences were observed in $I_{KV2,1}$ at various 235 voltages. In **panel 3C**, data points without asterisks are not considered significant. 236 Similarly, we measured the steady-state activation of the Kv1.5 current (I_{Kv1.5}) as shown 237 in **panel 3D** where male and female experimental data in myocytes are shown with blue 238 and black symbols. Properties of IKv1.5 steady-state activation G/Gmax show minimal sex-239 specific differences. The current-voltage relationship of IKv1.5 is shown from 240 measurements in female (blue, n=10) and male (black, n=7) myocytes in **panel 3E**. 241 Finally, the current-voltage relationship of the contribution from $I_{Ky1.5}$ and $I_{Ky2.1}$ to the total 242 voltage-gated current (IKVTOT) is shown in **panel 3F** with male and female data shown with black and blue symbols, respectively. Data points in panel D, panel E, and panel F 243 244 without asterisks are not significant. The table in panel 3H summarizes the sexdependent maximal conductance and the current response at specific voltages of -50 mV, 245 246 -40 mV, -30 mV, and -20 mV for both $I_{Kv1.5}$ and $I_{Kv2.1}$.

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To understand the contribution of each K⁺ current to the total voltage-gated current (I_{KVTOT}) in mesenteric vascular smooth muscle cells we built and optimized a Hodgkin-Huxley model to the data described above. First, we developed a model to describe the Kv2.1 current. The optimized model to Kv2.1 experimental data contains only a voltagedependent activation gating variable (X_{2.1act}). Since inactivation time is slow and is well estimated by steady-state³, we did not consider its effects in our model. The model of I_{Kv2.1} is described by:

255 (2) $I_{Kv2.1} = G_{Kv2.1} * X_{Kv2.1act} * (V - E_K)$

256

Where $G_{K2.1}$ is the maximal conductance of Kv2.1 channels and E_K is the Nernst potential for potassium. Parameters were optimized to male and female experimental data as shown for activation curves in **Figure 3A**. Model optimization to male and female time constants of activation (Kv2.1 TActivation) are shown as solid lines in **Figure 3B**. The model was also optimized to the I_{Kv2.1} current-voltage (I-V) relationships shown as solid lines in **Figure 3C**.

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Similarly, we developed a model for $K_V 1.5$. The model was optimized to the $K_V 1.5$ experimental data and contains only a voltage-dependent activation gating variable $(X_{Kv1.5act})$. The model of $I_{Kv1.5}$ is described by:

267

268 (3)
$$I_{KV1.5} = G_{Kv1.5} * X_{Kv1.5act} * (V - E_K)$$

269

270 $G_{K1.5}$ is the maximal conductance of Kv1.5 channels and E_K is the Nernst potential for 271 potassium. Parameters were optimized to male and female experimental data as shown 272 for activation curves in **Figure 3D**. The model was also optimized to the $I_{Ky1.5}$ current-273 voltage (I-V) relationships shown as solid lines in Figure 3E. From experiments, we 274 optimized the model to reproduce the overall time traces of Kv currents. The model 275 predicted that male and female myocytes have comparable time constants of activation 276 in $I_{Kv1.5}$ as shown in **Figure 3G.** Finally, the optimized model of the total voltage-gated 277 current (I_{KVTOT}) is shown in **Figure 3F**. The total voltage-gated K⁺ current (I_{KVTOT}) is the 278 sum of $I_{KV1.5}$ and $I_{KV2.1}$ mathematically described as:

279

280 (4) IKVTOT = IKV2.1 + IKV1.5

281

Notably, the main specific sex-specific differences observed in the total voltage-gated K⁺
 current (I_{KvTOT}) is attributable to the sex-specific differences in the current produced by
 K_v2.1 channels.

285

We next analyzed the contribution of large-conductance calcium-activated potassium (BK_{Ca}) channels to vascular smooth muscle cell electrophysiology. BK_{Ca} channels are

activated by membrane depolarization or increased [Ca²⁺]; and are expressed in the 288 289 membrane of vascular smooth muscle cells with α and β 1 subunits^{22,54,55}. In smooth 290 muscle cells, Ca²⁺ sparks are the physiological activators of BK_{ca} channels. We relied on 291 the assumption by Tong *et al.*⁵⁶ that BK_{Ca} currents (I_{BKCa}) are produced by two current 292 subtypes, one consisting of α subunits (I_{BK α}) and the other consisting of α and β 1 subunits 293 (I_{BKaβ1}). Experimental evidence indicates that BK_{Ca} channels with $\alpha\beta1$ subunits form 294 clusters in the plasma membrane in specialized junctional domains formed by the 295 sarcoplasmic reticulum and the sarcolemma. BK_{Ca} channels with $\alpha\beta1$ subunits colocalize 296 with ryanodine receptors (RyRs) to in the junctional domains. During a Ca²⁺ spark, $[Ca^{2+}]_i$ elevations ranging from 10 to 100 µM activate BK_{Ca} channels^{38,39,52,57,58}. In our model, 297 298 Ca²⁺ sparks are the physiological activators of BK_{Ca} channels.

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300 The mathematical formulation of the BK_{Ca} with $\alpha\beta1$ current (I_{BK\alpha\beta1}) was optimized to fit the experimental whole-cell electrophysiological data from Bao and Cox⁵⁴ obtained at room 301 302 temperature with a BK_{ca} channel α subunit clone from mSlo-mbr5 and a β 1 subunit clone from bovine expressed in *Xenopus laevis* oocytes⁵⁴. Experimental data for steady-state 303 304 activation and time constants of activation are shown in Panel 4A and Panel 4B 305 respectively. The activation gating variable (X_{ab}) depends on both voltage and junctional calcium ([Ca²⁺]_{Jun}). The activation gate was adapted from the Tong-Taggart model⁵⁶. The 306 307 model of $I_{BK\alpha\beta1}$ is described by:

308

309 (5)
$$I_{BK\alpha\beta1} = P_{BKca} * X_{ab}(V, [Ca]_{Jun}) * \frac{z_K^2 F^2 V}{RT} \left(\frac{[K]_{in} e^{\frac{Z_K F V}{RT}} - [K]_{out}}{e^{\frac{Z_K F V}{RT}} - 1} \right)$$

310

Where P_{BKCa} is the BK_{Ca} ion permeability, R is the gas constant, F is Faraday's constant, and z_{K} is the valance of the potassium ions. Model optimization to activation curves are shown with solid lines in **Figure 4A** at three different $[Ca^{2+}]_{Jun}$ concentrations 1 μ M, 10 μ M, and 100 μ M. The results from the steady-state activation measurements at 10 μ M are also in agreement with the experimental data in vascular myocytes in *bufo marinus*⁵⁸ (green symbols) which suggests that BK_{Ca} channels are exposed to a mean junctional Ca²⁺ concentration ($[Ca^{2+}]_{Jun}$) of 10 μ M. Time constants of activation were measured

experimentally at $[Ca^{2+}]_{Jun} = 0.003 \mu M$, our model was optimized and fit under the same 318 319 conditions shown in Figure 4B as solid lines. Notably when the model was run under 320 predicted $[Ca^{2+}]_{Jun} = 10 \ \mu M$ conditions as shown in **Figure 4B** dashed lines, there was no 321 effect of the change in $[Ca^{2+}]_{Jun}$ on the time constant. The predicted current-voltage (I-V) 322 relationships of $I_{BK\alpha\beta1}$ are shown in **Figure 4C** using three different $[Ca^{2+}]_{Jun}$ 323 concentrations; 1 µM, 10 µM, and 100 µM. We observed that the I-V curves are similar at 324 $[Ca^{2+}]_{Jun}$ concentrations of 10 μ M (black trace) and 100 μ M (orange trace) but markedly reduced when $[Ca^{2+}]_{Jun} = 1 \mu M$ (blue trace). As expected, the amplitude of the current 325 326 shown in the I-V curves in Figure 4D is sensitively dependent on the number of BKca channels as shown, we set [Ca²⁺]_{Jun}= 10 µM and simulated the I-V curves using a BK_{Ca} 327 328 cluster size of 4, 6, 8 and 10 channels.

329

330 In vascular smooth muscle cells, the membrane potential over the physiological range of 331 intravascular pressures is less negative than the equilibrium potential of potassium ($E_{K} =$ 332 -84 mV), suggesting active participation of inward currents regulated by sodium conductance^{19,59,60}. It has been postulated that basally activating TRP channels generate 333 334 nonselective cations currents (I_{NSC}) that depolarize the membrane potential. We built a model for I_{NSC} as linear and time-independent cation current permeable to K⁺ and Na⁺ 335 336 with permeability ratios P_{Na} : P_{K} =0.9:1.3 adapted from Tong-Taggart model with a 337 reversal potential (ENSC) described by:

338

339 (6)
$$E_{NSC} = \left(R * \frac{T}{F}\right) * log\left(\frac{P_K * K_{out} + P_{Na} * Na_{out}}{P_K * K_{in} + P_{Na} * Na_{in}}\right);$$

340

where R is the gas constant, F is the Faraday's constant, T is the temperature, Na_{in} and
K_{in} are the intracellular sodium and potassium intracellular concentrations. Similarly,
Na_{out} and Na_{out} are the extracellular sodium and potassium concentrations. The model
of I_{NSC} is described by

- 345
- 346 (7) $I_{NSC} = I_{NaNSC} + I_{KNSC}$
- 347 (8) $I_{NaNSC} = G_{NaNSC} * (V E_{NSC})$
- 348 (9) $I_{KNSC} = G_{KNSC} * (V E_{NSC})$

13

349	
350	Where I_{NaNSC} represents sodium current contribution, I_{KNSC} represents potassium current
351	contribution, and GNANSC and GKNSC are the maximal conductances of the contributing
352	sodium and potassium currents. In addition, we also included models for leak currents of
353	ion i calculated as
354	
355	(10) $I_{i,b} = G_{i,b} (V - E_i)$
356	
357	Where the Nernst potential of ion i with valance z_i is given by:
358	
359	(11) $E_i = RT/z_iF \ln ([i]_{out}/[i]_{in}), i = Na^+, K^+ and Ca^{2+}$
360	
361	Where R is the gas constant, F is the Faraday's constant, T is the temperature and $[i]_{out}$
362	denotes the extracellular concentration of ion i. FF
363	
364	The remaining ionic currents, pumps, and transporters were optimized to data available
365	in the experimental literature and/or taken from computational models of vascular smooth
366	muscle and cardiac cells. The sodium-potassium pump (I_{NaK}) current was modeled using
367	data from smooth muscle cells from mesenteric resistance arteries of the guinea pig ^{56,61}
368	and the voltage dependency was adapted from the Luo-Rudy II model ⁴¹ . The sodium-
369	calcium exchanger current (I_{NCX}) was adapted from the formulation in the ten Tusscher
370	model ⁶² and the Luo-Rudy II model ⁴¹ . Finally, the sarcolemma calcium pump (IPMCA)
371	current was adapted from the Kargacin model ⁶³ .
372	

373 (12)
$$N_{pow} = \left(Q10^{\frac{T-309.2}{10}}\right);$$

374 (13)
$$N_1 = \frac{(K_{out}^{1.1})}{K_{out}^{1.1} + KmNaK_K^{1.1}};$$

375 (14)
$$N_2 = \frac{(Na_{in}^{1.7})}{Na_{in}^{1.7} + KmNaK_{Na}^{1.7}};$$

376 (15) $N_0 = \frac{1.0}{1 + (0.1245 * exp(-0.1 * V * \frac{F}{R * T})) + (2.19e - 3 * (e^{(\frac{Na_{out}}{49.71})}) * e^{(-1.9 * V * \frac{F}{R * T})})};$

14

377 (16)
$$I_{NaK} = I_{NaK_{max}} * N_1 * N_2 * N_0 * N_{pow}$$

378 (17)
$$phi_F = exp\left(gammax * V * \frac{F}{R*T}\right)$$

379 (18)
$$phi_R = exp\left((gammax - 1) * V * \frac{F}{R*T}\right)$$

380 (19)
$$X_{NCX} = \frac{(Na_{in}^3) * Ca_{out} * phi_F - (Na_{out}^3) * Ca_{i*} phi_R}{1 + 0.0003 * ((Na_{out}^3) Ca_{in} + (Na_{in}^3) * Ca_{out})};$$

$$381 \quad (20) \quad I_{NCX} = P_{NCX} * X_{NCX}$$

382 (21)
$$I_{PMCA} = I_{PMCAbar} * \frac{Ca_i^2}{Ca_i^2 + K_{mPMCA}^2}$$

383

We next set out to connect the ionic models and models of Ca²⁺ handling to make 384 predictions in the whole cell. In **Figure 5**, experimental data indicate that the electrical 385 386 activity of isolated mesenteric smooth muscle cells in male and female myocytes recorded 387 in current-clamp mode, is characterized by an oscillating membrane potential under 388 physiological conditions. The membrane potential is marked by repetitive spontaneous 389 transient hyperpolarization (TH), a ubiquitous feature of vascular smooth muscle cells^{57,64–66} as shown in **panel 5A.** Both male (black trace) and female (blue trace) 390 391 myocytes exhibited membrane hyperpolarizing transients in the potential range of -50 to 392 -20 mV. Notably, we observed that female myocytes always maintained a higher 393 depolarizing state between the hyperpolarization events compared to male myocytes.

394

395 We assessed the predictive capacity of our *in silico* model by comparing it to experimental 396 data. We first compared the morphology of the membrane potential in experiments **panel** 397 5A versus simulations panel 5B in male and female myocytes. Upon comparative 398 analysis between male and female experimental data and simulations, we noted that the 399 baseline membrane potential for male myocytes was around -40 mV, while female 400 myocytes exhibited a slightly more depolarized membrane potential at approximately -30 401 mV. Despite these variations in baseline membrane potential, both male and female 402 myocytes presented similar peak hyperpolarization values of approximately 10-15 mV, 403 ranging from -50 mV to -30 mV. Similarly, the frequency of THs from multiple myocytes 404 was calculated to be 1 to 2.8 Hz in the range of -50 mV to -30 mV which is identical to 405 the simulated frequency.

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407 In the physiological range in which smooth muscle cells operate (-50 to -20 mV), ionic 408 currents are small and produced by the activation of a small number of ion channels. 409 Local fluctuations in the function of ion channels lead to noisy macroscopic signals that 410 are important to the variability of vascular smooth muscle cells¹⁹. In addition, smooth 411 muscle cells are subject to high input resistance where small perturbations can lead to large changes in the membrane potential^{19,29}. To approximate the physiological realities, 412 413 we applied two sources of noise to our deterministic in silico model to simulate the 414 stochastic fluctuations. The first source of the noise was introduced by adding a 415 fluctuating current term to the differential equations describing changes in membrane 416 potential (dV/dt) which represents the combined effect of the stochastic activity of ion 417 channels in the plasma membrane⁴⁶. Second, we introduced noise into the [Ca]_{SR} to 418 replicate the physiological responses consistent with those observed in experimental 419 studies⁴⁰. Simulated whole-cell membrane potential with physiological noise is shown in 420 Figure 5B in male (black trace) and female (blue trace) myocytes.

421

422 We conducted a sensitivity analysis to determine which model parameters could underlie 423 the sex-specific differences observed in the experimental data. It is important to note that 424 we have experimental data indicating the amplitude and kinetics for a variety of currents 425 in male and female myocytes. For this reason, those model components were fit to the 426 data, fixed, and were not subject to sensitivity analysis. Our analysis, which focused solely 427 on variations in maximal conductance and maximal ion transport rates of the 428 transmembrane currents, indicated that the non-selective cation currents (INSC) and 429 delayed rectifier currents ($I_{KVTOT} = I_{KV2.1} + I_{KV1.5}$) interact to regulate the baseline membrane 430 potential in both male and female vascular smooth muscle myocytes (**Figure 5C**). Given 431 that I_{KVTOT} responds to depolarization, the primary stimulus that triggers depolarization 432 was determined to be attributable solely to the non-selective cation currents (I_{NSC}). 433 Indeed, when we adjusted the conductance of the non-selective cation currents and 434 implemented an increase in the conductance of I_{NSC} in the female model, we readily 435 reproduced the sex-specific baseline membrane potential observed experimentally 436 (Figure 5A).

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438 Next, using the whole-cell vascular smooth muscle myocyte computational model, we 439 investigated the sex-specific differences in the contribution to total voltage-gated current 440 (I_{KVTOT}) in mesenteric vascular smooth muscle cells. An interesting prediction from the *in* 441 silico simulations is that at different depolarizing states (-45, -40, and -35 mV) induced by changing the conductance of nonselective cationic leak currents (I_{NSC}), the contribution 442 443 of IKv2.1 and IKv1.5 to IkvTOT is different based on sex. In male vascular myocytes, the 444 contribution to total voltage-gated current (I_{KVTOT}) is largely attributable to the current 445 produced by $K_{V1.5}$ channels as shown in the lower panel in **Figure 6A**. Our results are consistent with previous studies^{35,67,68} in animal rodent male models showing the 446 447 characteristic behavior of I_{Kv1.5} to control membrane potential. However, the model 448 predicts that in female myocytes, the contribution to total voltage-gated current (IKVTOT) is 449 largely provided by the current produced by Kv2.1 channels as shown in the upper panel 450 in Figure 6B. To illustrate this point quantitatively, at a membrane potential of -40 mV, 451 the contribution of IKVTOT from IKV1.5 and IKV2.1 86% and 14%, respectively, in male 452 myocytes compared to female myocytes in which the contribution from $I_{Ky1.5}$ and $I_{Ky2.1}$ is 23% and 77%, respectively. Regardless of the depolarization state at -45, -40, or -35 mV, 453 454 the profiles for male and female myocytes remain essentially the same as shown in 455 Figures 6C, 6D, and 6E. The in silico simulations suggest a distinctive sex-based 456 function of Kv1.5 and Kv2.1 channels that produce the delayed rectifier currents to 457 counterbalance the effect of inward currents causing graded membrane potential 458 depolarizations.

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460 Having explored the regulation of graded membrane potential by the activation of IKVTOT 461 to counterbalance the nonselective cations currents (I_{NSC}), we next explored the effects 462 of steady membrane depolarization in the in silico vascular smooth muscle cell myocyte 463 model on I_{Ca} in male and female myocytes. We predicted I_{Ca} in our male and female 464 simulations at steady-state membrane depolarization after simulation for 500 seconds. 465 We observed that as the membrane depolarizes from -55 to -35 mV, I_{ca} in male myocytes increased from 0 to 1.0 pA while in female myocytes I_{Ca} increased from 0 to 1.5 pA as 466 467 shown in **Figure 7A**, suggesting that I_{Ca} is larger in female compared to those of male

myocytes. We recorded the predicted $[Ca^{2+}]_i$ and observed that I_{Ca} led to a higher calcium 468 469 influx in female compared to male simulations as shown in **Figure 7B**. To illustrate in 470 detail, we show in **Figure 7C-D**, time traces of *in silico* predictions of membrane voltage 471 at -40 mV (top panel), I_{Ca} (middle panel), and [Ca²⁺]_i (lower panel) corresponding to the 472 male and female data points indicated by black and blue arrows respectively shown in 473 Figure 7A-B. In the male case (Figure 7C), at a steady membrane potential of -40 mV, 474 L-type calcium Cav1.2 channels produced a current of 0.5 pA. However, in female 475 simulations (Figure 7D), we observed that at a steady membrane potential of -40 mV, L-476 type calcium Cav1.2 channels produced a current of 0.65 pA. We calculated that at -40 477 mV, two Ca $\sqrt{1.2}$ channels are needed to sustain 0.5 pA of current in male myocytes while 478 three Cav1.2 channels are needed to sustain 0.65 pA of current in female myocytes. 479 Although the sex-specific differences in male and female simulations at -40 mV are small, a 15 nM difference in the overall response of [Ca²⁺]; can have a profound effect on the 480 481 constriction state of the myocytes. The predictions from the Hernandez-Hernandez model 482 provide a comprehensive picture of physiological conditions and support the idea that a small number of Ca_V1.2 channels supply the steady Ca²⁺ influx needed to support a 483 maintained constricted state in small arteries and arterioles^{53,69}. The differences between 484 485 males and females are notable in the context of observations indicating varied sex-based responses to antihypertensive agents that target the Ca²⁺ handling system in vascular 486 487 smooth muscle cells.

488

489 Next, we simulated the effects of calcium channel blocker nifedipine on Ica at a steady 490 membrane potential of -40 mV in male and female simulations. Briefly, previous studies⁷⁰ 491 have shown that at the therapeutic dose of nifedipine (i.e., about 0.1 µM) L-type Cav1.2 492 channel currents are reduced by about 60-70%. Accordingly, we decreased Ica in our 493 mathematical simulations by the same extent. In Figure 7C-D, we show the predicted 494 male (grav) and female (pink) time course of membrane voltage at -40 mV (top panel). 495 Ica (*middle panel*), and [Ca²⁺]; (*lower panel*). First, we observed that in both male and 496 females 0.1 µM nifedipine modifies the frequency of oscillation in the membrane potential, 497 by causing a reduction in oscillation frequency. Second, both male and female simulations 498 (middle panels) show that 0.1 μ M nifedipine caused a reduction of I_{Ca} to levels that are

499 very similar in male and female myocytes following treatment. Consequently, the 500 reduction of I_{Ca} causes both male and female simulations to reach a very similar baseline 501 $[Ca^{2+}]_i$ of about 85 nM (lower panels). As a result, simulations provide evidence supporting 502 the idea that $Ca_V 1.2$ channels are the predominant regulators of intracellular $[Ca^{2+}]$ entry 503 in the physiological range from -40 mV to -20 mV. Importantly, these predictions also 504 suggest that clinically relevant concentrations of nifedipine cause larger overall reductions 505 in Ca^{2+} influx in female than in male arterial myocytes.

506

507 Thus far, we have shown the development and application of models of vascular smooth 508 muscle myocytes incorporating measured sex-specific differences in currents from male 509 and female isolated cells. Given that hypertension is essentially a consequence of the 510 spatial organization and function of smooth muscle cells^{71,72}, we next expanded our study 511 to include a one-dimensional (1D) tissue representation of electrotonically coupled tissue 512 by connecting arterial myocytes in series.

513

514 A well-known phenomenon in excitable systems is that electrotonic coupling between 515 cells results in the minimization of individual cellular differences, thereby producing a smoothing effect across the tissue^{73–75}. We simulated 400 female or 400 male vascular 516 517 smooth muscle myocytes and set the gap junctional conductivity to zero to uncouple the 518 simulated cells. As expected, the uncoupled cells in both male and female cases 519 demonstrated the characteristic behavior of arterial myocytes, exhibiting spontaneous hyperpolarization. Of the 400 cells, we show the simulated representative traces of Cell 520 521 1, Cell 50, and Cell 100 for female (Figure 8A) and male (Figure 8B).

522

Next, we modeled 400 cells but with electrotonic coupling by setting the simulated gap junctional resistance to 71.4 Ω cm² ⁷⁶. In this case, we observed that the spontaneous hyperpolarizations, previously observed in the uncoupled cells, diminished when cells were coupled. The overall smoothing effect observed in **Figure 8C** is attributed to the electrotonic coupling and consequential influence of neighboring cells. The electrical response is consistent across the spatial domain for both male (**Figure 8C**; *black trace*) and female (**Figure 8C**-*blue trace*) one-dimensional tissue representations. Notably, the

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model predicts a more depolarized female membrane potential in the one-dimensional
tissue representations consistent with experimental measurements as shown in Figure
8D.

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534 Having developed an idealized model of a vessel, we set out to validate the model 535 predictions of variable [Ca²⁺]; between males and females by comparing the computed 536 calcium signaling in vascular smooth muscle with experimental recordings O'Dwyer et 537 al.²⁰. Given that membrane potential predominantly governs calcium influx in vascular 538 smooth muscle¹⁴, we varied the conductance of the nonselective cation currents (I_{NSCC}) in our simulations. Tuning of INSCC was performed to replicate the effects of pressure-539 540 induced membrane depolarization, which results in activation of the voltage-gated L-type Ca²⁺ channels and increases [Ca²⁺]_i. 541

542

543 Our simulations (lines) are well validated by experimental recordings (symbols) in **Figure** 544 **9A**. A distinctive feature from the model prediction, which was validated by experimental 545 recordings is the observation that female (**Figure 9A**, blue trace and symbols) vessels 546 accommodate more $[Ca^{2+}]_i$ compared to male (**Figure 9A**, black trace and symbols) 547 vessels. Intriguingly, the mechanism of different $[Ca^{2+}]_i$ in male and female vessels was 548 revealed in single-cell simulations, which showed attributable sex-based differences in L-549 type Ca^{2+} currents.

550

Finally, in our simulations, we computed the effects of [Ca²⁺], after the application of 551 552 clinically relevant calcium channel blocker nifedipine. We observed a substantial 553 reduction of [Ca²⁺]_i in both male (Figure 9A, dashed black line) and female (Figure 9A, 554 dashed blue line). Significant differences were found in the physiological range of 555 intravascular pressure from 40 to 120 mmHg. In the summary data (Figure 9B), we 556 quantified the relative change of [Ca²⁺]; in male (*black*) and female (*blue*) after the 557 application of 0.1 mM L-type Ca²⁺ channel blocker nifedipine at 80 mmHg and 120 mmHg. Our results show that nifedipine, when applied to male vessels, decreases $[Ca^{2+}]_i$ by 22% 558 559 and 25% at 80 mmHg and 120 mmHg, respectively. However, the same dose of nifedipine when applied to female vessels decreases $[Ca^{2+}]_i$ by 38% and 45% at 80 mmHg and 120 560

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561 mmHg. The results suggest that female arterial smooth muscle is more sensitive to 562 clinically used Ca²⁺ channel blockers than male smooth muscle.

563

564 **Discussion**

565 Here, we describe the development, validation, and application of an *in-silico* model to simulate and understand the mechanisms of electrical activity and Ca²⁺ dynamics in a 566 567 single mesenteric vascular smooth muscle cell. The Hernandez-Hernandez model is the first model to incorporate sex-specific differences in voltage-gated Ky2.1 and Cay1.2 568 channels and predicts sex-specific differences in membrane potential and Ca²⁺ signaling 569 570 regulation in the smooth muscle of both sexes from systemic arteries. In the pursuit of 571 stratifying sex-specific responses to antihypertensive drugs, we expanded our exploration 572 to encompass a one-dimensional (1D) tissue representation. Such an approach allowed us to simulate and predict the impact of Ca²⁺ channel blockers within a mesenteric vessel. 573 574 Notably, the computational framework can be expanded to forecast the impact of 575 antihypertensives and other perturbations from single-cell to tissue-level simulations.

576

To specifically investigate the impact of sex-specific differences measured from ion channel experiments and their impact on membrane potential and $[Ca^{2+}]_i$, we focused on the isolated myocyte in the absence of complex signaling pathways. We first explored the effects of Cav1.2 and Kv2.1 channels on membrane potential as experimental data suggest key sex-specific differences in channel expression and kinetics. Notably, the peak of the current-voltage (I-V) relationship of L-type Cav1.2 current is 40% smaller in male compared to female myocytes (**Figure 2D**).

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Similarly, the peak current-voltage (I-V) relationship of the voltage-gated Kv2.1 current ($I_{Kv2.1}$) is 70% smaller in male compared to female myocytes at +40 mV (**Figure 3C**). O'Dwyer and coauthors²⁰ showed sex-dependent expression of Kv2.1 in the plasma membrane, where male arterial myocytes have a total of about 75,000 channels compared to 183,000 channels in female myocytes. Notably, less than 0.01% of channels are conducting in male and female myocytes. In the computational model, we found that to reproduce the experimentally measured amplitude of the Kv2.1 I-V curve (**Figure 3C**),

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a maximum of ~44 male Kv2.1 channels was sufficient to reproduce the peak current (68.8 pA at 40 mV). In contrast, ~143 channels were predicted to be needed in female myocytes to reproduce the experimentally measured peak current (226.42 pA at +40 mV) of the Kv2.1 I-V relationship. Modeling and simulation led to the prediction that in male arterial myocytes, I_{KvTOT} is largely dictated by Kv1.5 channels. In contrast, in female arterial myocytes, Kv2.1 channels dominate I_{KvTOT} (Figure 3F).

598

An important aspect of the Hernandez-Hernandez model is that it includes Ca²⁺-mediated 599 600 signaling between RyRs in the junctional SR and BK_{Ca} channel clusters in the nearby 601 sarcolemma membrane. This section of the model is similar to the one included in the 602 Karlin model⁵ with some modifications. The Karlin model described how subcellular junctional spaces influence membrane potential and [Ca²⁺]; in response to intravascular 603 604 pressure, vasoconstrictors, and vasodilators. In this study, we reduced the complexity of the model representation of subcellular Ca²⁺ signaling spaces to include just three 605 606 compartments: the cytosol, SR, and the SR-sarcolemma junction. Our model represents 607 on average, the behavior of a single junctional SR unit that is functional in a cell at a time. The model uses a deterministic approach but mimics the process of production of Ca²⁺ 608 sparks that activate BK_{Ca} channel clusters²⁹. We represented the activity of the RyRs in 609 the junctional domain deterministically in the model so that Ca²⁺ spark- BK_{Ca} currents 610 611 occur at a frequency of about 1 Hz at -40 mV in a space equivalent to 1% of the total cell surface area of the plasma membrane²². 612

613

Based on experimental observations, the Hernandez-Hernandez computational model employs three key assumptions: First, Ca^{2+} sparks in the junctional domain are initiated by activation of RyRs, where RyR gating opening probability is correlated with SR load. Second, Ca^{2+} sparks lead to a $[Ca^{2+}]_{Jun}$ increase between 10-20 mM to match the amplitude measured in experiments (**Figure 4A**)^{54,58}. Third, activation of BK_{Ca} channels and the resultant current amplitude derives from the experimentally observed spontaneous outward currents (STOCs) in both amplitude and morphology.

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622 Notably, model simulations revealed important mechanisms that may underlie 623 experimental observations in measurements of membrane potential (Figure 5A). The 624 model predicts that the mechanism of intrinsic oscillatory behavior in the vascular 625 myocytes results from a delicate balance of currents. Activation of non-selective cation currents (I_{NSC}) likely causes membrane depolarization, but the delayed rectifier currents 626 627 (I_{KVTOT}) oppose them, resulting in membrane potential baseline in the physiological range of -45 to -20 mV. Interestingly, the voltage-gated L-type Cav1.2 currents activation 628 629 threshold sits within this range at ~-45 mV. Therefore, small increases in INSC can 630 overwhelm IKVTOT below -20 mV and result in sufficient depolarization to bring the 631 membrane potential to the threshold for activation of I_{Ca} . It is important to note that I_{KVTOT} 632 increases sharply upon depolarization from -45 to -20 mV, resulting in tight control of 633 membrane potential and prevention of large transient depolarization resulting from INSC. Activation of L-type Ca²⁺ channels upon depolarization and subsequent Ca²⁺ release 634 635 within the small volume junction then activates the BK_{ca} channels, which results in 636 hyperpolarization. Hyperpolarization reignites the oscillatory cascade as an intrinsic 637 resetting mechanism. Since vascular myocytes are subject to substantial noise from the 638 stochastic opening of ion channels in the plasma membrane, and fluctuations in the local 639 junctional domain components, such as the SR load, RyR opening, and BK_{ca} channel 640 activity, we included noise in the simulation. To simulate the physiological noise in the 641 vascular smooth muscle cell (Figure 5B), we added Gaussian noise to the dV/dt 642 equations and [Ca]_{SR}.

643

644 Female mesenteric artery myocytes are more depolarized than male myocytes at physiological intravascular pressures²⁰. Our model suggests that female myocytes are 645 646 more depolarized than male myocytes due to larger non-selective cation currents in 647 female compared to male myocytes, most likely due to the activation of Na⁺-permeable 648 TRP channels. To our knowledge, the only TRP channels found to regulate the 649 membrane potential of mesenteric artery smooth muscle are TRPP1 and TRPP2^{11,12}. Future work will have to determine if TRPC6⁸ and TRPM4^{9,10}, which have been shown to 650 651 mediate the myogenic response of cerebral artery smooth muscle, and/or other non-652 selective cation channels also depolarize mesenteric artery smooth muscle⁷⁷.

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653

The Hernandez-Hernandez model predicts that very few channels (based on total current amplitude) are likely to control the baseline fluctuations in membrane potential in the physiological range of -60 to -20 mV. The intrinsic oscillatory properties of the vascular myocyte operating in the low voltage regime under conditions of high resistance membrane are similar to other types of oscillatory electrical cells including cardiac pacemaker cells.

660

661 As shown in (Figure 6), the model predicts that, at -40 mV, the amplitude of steady-state 662 Ky2.1 currents is about 0.8 pA in male and 3.3 pA in female arterial myocytes, indicating that the contribution of Ky2.1 and Ky1.5 channels to membrane potential is different in 663 664 males and females. At -30 mV, it is 2.34 pA and 9.2 pA in male and female myocytes respectively. Assuming a single channel current at -40 and -30 mV of 0.7 pA, we 665 666 calculated that, on average, in male myocytes a single channel is open at -40 mV and 3 667 channels are open at any particular time at -30 mV. In female myocytes, 6 channels are 668 predicted to be open at -40 mV, while 13 are predicted to be active at -30 mV.

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670 The Hernandez-Hernandez model also allowed us to calculate the number of Ca_V1.2 671 channels needed to sustain the steady-state concentration of [Ca²⁺], in the physiological 672 range from -60 to -20 mV (Figure 7). The model predicts that at -40 mV in mouse male 673 myocytes, 2 channels were required to generate 0.5 pA of steady-state Ca_V1.2 current. 674 On the other hand, we found that in female myocytes, 3 channels were sufficient to 675 generate 0.65 pA of Ca_V1.2 current. These data are consistent with the work of Rubart et 676 al.⁶⁹, which suggested that steady-state Ca²⁺ currents at -40 mV were likely produced by 677 the opening of 2 Ca \vee 1.2 channels in rat cerebral artery smooth muscle cells.

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The observation that a very small number of the conducting Kv2.1 and Cav1.2 channels are involved in the regulation of membrane potential and Ca²⁺ influx in male and female arterial myocytes at physiological membrane potentials is important for several reasons. *First*, the analysis suggests that small differences in the number of Kv2.1 and Cav1.2 channels can translate into large, functionally important differences in membrane

potential and $[Ca^{2+}]_i$ and hence affect and control myogenic tone under physiological and pathological conditions. *Second*, the small number of K_V2.1 and Ca_V1.2 channels gating between -40 and -30 mV likely makes smooth muscle cells more susceptible to stochastic fluctuations in the number and open probabilities of these channels than in cells where a large number of channels regulate membrane excitability and Ca²⁺ influx (e.g., adult ventricular myocyte⁷⁸). This, at least in part, likely contributes to Ca²⁺ signaling heterogeneity in vascular smooth muscle.

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Hypertension fundamentally manifests through the spatial organization of cellular 692 693 components, particularly evident in the context of the tunica media, the middle layer of 694 vessels is predominantly constituted of smooth muscle cells which play a pivotal role in vessel contraction and relaxation^{71,72}. Such intricate biological machinery is imperative in 695 orchestrating the regulation of blood flow and blood pressure. Our approach began with 696 697 a process of distillation, aiming to shed light on cellular mechanisms within isolated 698 vascular myocytes from small systemic vessels and arterioles, which control blood 699 pressure, of both male and female mice.

700

701 Earlier research has confirmed that in mesenteric arteries, the pathogenesis leading to hypertension is largely determined by the downregulation of Kv2.1³⁶ and/or Kv1.5^{67,79} and 702 a concurrent increase in the activity of Cav1.2⁸⁰ channels. Building upon this knowledge. 703 704 we broadened our study to encompass a one-dimensional (1D) tissue model of 705 electrotonically linked tissue, achieved by connecting arterial myocytes in series. The 1D 706 cable model has anatomical relevance because the structure of third and fourth-order 707 mesenteric arteries have a singular layer of vascular myocytes encircling the lumen in a 708 cylindrical arrangement. The cable structure is analogous to an "unrolled" or lateral 709 arrangement of the vessel. Such an approach allowed a conceptual framework to bridge 710 the gap between understanding the combined effects of membrane potential and $[Ca^{2+}]_i$ 711 in isolated cells and in the wider context of small vessels.

712

For instance, previous studies have proposed that gap junctions enable vessels to function in a way that is analogous to a large capacitor^{57,81}. The gap junctions actively

715 filter and transform single-cell electrical activity into sustained responses across the 716 tissue⁸¹. Recent studies add to this understanding by demonstrating that Connexin 37 717 (Cx37), a component of these gap junctions, seems to be expressed in the mesenteric 718 arteries⁸². In our simulations, we showed (Figure 8A-B) that indeed uncoupled cells 719 exhibit a spontaneous oscillatory behavior which studies have confirmed is not an artifact 720 due to isolation from the vessel but rather an intrinsic behavior required to sustain 721 electrical signals. When the cells are connected (Figure 8C) the spontaneous 722 hyperpolarization previously observed in the uncoupled cells diminished, the effect is 723 attributed to the electrotonic coupling and consequential influence of neighboring cells. In 724 addition, in our simulations, we found that it is required to have stochastic fluctuations to 725 allow the system to average the membrane potential behavior that dictates the amount of 726 [Ca²⁺]_i in the vessels.

727

Regarding, Cav1.2 channels, simulations forecast the clinically relevant concentrations (0.1 μ M) at which common Ca²⁺ channel blockers (e.g., nifedipine) effectively block Cav1.2 channels in both male and female smooth muscle (**Figure 9**). Our simulations in isolated arterial myocytes and in the one-dimensional (1D) tissue model suggest heightened sensitivity to calcium channel blockers in the female compared to male.

733

734 The model predictions are aligned with documented sex-specific differences in 735 antihypertensive drug responses^{83,84}. Previous studies, notably by Kloner et al., have 736 illustrated this point quantitatively, highlighting a more pronounced diastolic BP response 737 in women (91.4%) compared to men (83%) when treated with dihydropyridine-type 738 channel blockers, such as amlodipine. Importantly, this distinction persisted even after 739 adjusting for confounding factors such as baseline BP, age, weight, and dosage per 740 kilogram⁸⁴. Another interesting observation from Kajiwara et al. emphasizes that 741 vasodilation-related adverse symptoms occur more frequently in younger women (<50 742 years) compared to their male counterparts, again suggesting a heightened sensitivity to 743 dihydropyridine-type calcium channel blockers⁸⁵.

744

745 Limitations

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746

747 The model presented here describes the necessary and sufficient ion channels, pumps, 748 and transporters to describe the electrical activity and Ca²⁺ signaling of an isolated 749 mesenteric smooth muscle cell in the absence of complex signaling pathways. Such an 750 approach enabled us to perform a component dissection to analyze the sex-specific 751 differences observed in the fundamental electrophysiology of male and female myocytes. 752 However, it is well known that vascular smooth muscle cells are subject to a plethora of 753 stimuli from endothelial cells, neurotransmitters, endocrine, and paracrine signals⁵. The 754 next phase of the project includes an expansion of the model to incorporate receptor 755 mediated signaling pathways that are essential for blood pressure control.

756

Excitation-contraction coupling refers to an electrical stimulus that drives the release of calcium from the sarcoplasmic reticulum and results in the physical translocation of fibers that underlies muscle contraction. In the present model, we did not explicitly consider the mechanical description of muscle contraction. Nevertheless, we can imply contractile effects by tracking membrane potential and the elevation of $[Ca^{2+}]_i$ as a proxy.

762

To conclude, we developed and present the Hernandez-Hernandez model of male and female isolated mesenteric vascular myocytes. An additional limitation of our study is the reliance on predominantly murine data. Although mouse arteries do present numerous parallels with human arteries—including analogous intravascular pressure-myogenic tone relationships, resting membrane potentials, and the expression of typical ionic channels like Ca_V1.2, BKCa channels, and RyRs^{86–88}. Future research should assess the direct applicability and implication of our findings in human subjects.

770

771 **Conclusions**

The Hernandez-Hernandez model of the isolated mesenteric vascular myocyte was informed and validated with experimental data from male and female vascular myocytes. We then used the model to reveal sex-specific mechanisms of Kv2.1 and Cav1.2 channels in controlling membrane potential and Ca²⁺ dynamics. In doing so, we predicted that very few channels are needed to contribute to and sustain the oscillatory behavior of the

777 membrane potential and calcium signaling. We expanded our computational framework 778 to include a one-dimensional (1D) tissue representation, providing a basis for simulating 779 the effect of drug effects within a vessel. The model predictions suggested differences in 780 the response of male and female myocytes to drugs and the underlying mechanisms for 781 those differences. These predictions may constitute the first step towards better 782 hypertensive therapy for males and females.

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786 MATERIALS AND METHODS

- 787 Section 1. Experimental
- 788

789 **1.1 Animals**

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791 This study was performed in strict accordance with the recommendations in the Guide for 792 the Care and Use of Laboratory Animals of the National Institutes of Health. All of the 793 animals were handled according to approved institutional animal care and use committee 794 (IACUC) protocols of the University of California Davis. IACUC protocol number is 22503. 795 8- to 12-week-old male and female mice C57BL/6J (The Jackson Laboratory, 796 Sacramento, CA) were used in this study. Animals were housed under standard light-dark 797 cycles and allowed to feed and drink ad libitum. Animals were euthanized with a single lethal dose of sodium pentobarbital (250 mg/kg) intraperitoneally. All experiments were 798 799 conducted in accordance with the University of California Institutional Animal Care and 800 Use Committee guidelines.

801

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808

809 **1.2 Isolation of arterial myocytes from systemic resistance arterioles**

Third and fourth-order mesenteric arteries were carefully cleaned of surrounding adipose and connective tissues, dissected, and held in ice-cold dissecting solution (Mg²⁺-PSS; 5 mM KCl, 140 mM NaCl, 2mM MgCl₂, 10 mM glucose, and 10 mM HEPES adjusted to pH 7.4 with NaOH). Arteries were first placed in dissecting solution supplemented with 1.23 mg/ml papain (Worthington Biochemical, Lakewood, NJ) and 1 mg/ml DTT for 14 minutes at 37°C. This was followed by a second 5-minute incubation in dissecting solution supplemented with 1.6 mg/ml collagenase H (Sigma-Aldrich, St. Louis, MO), 0.5 mg/ml

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elastase (Worthington Biochemical, Lakewood, NJ), and 1 mg/ml trypsin inhibitor from *Glycine max* (Sigma-Aldrich, St. Louis, MO) at 37°C. Arteries were rinsed three times with
dissection solution and single cells were obtained by gentle trituration with a wide-bore

- glass pipette. Myocytes were maintained at 4°C in dissecting solution until used.
- 821

822 **1.3 Patch-clamp electrophysiology**

All electrophysiological recordings were acquired at room temperature (22–25°C) with an Axopatch 200B amplifier and Digidata 1440 digitizer (Molecular Devices, Sunnyvale, CA). Borosilicate patch pipettes were pulled and polished to resistances of 3-6 M Ω for all experiments using a micropipette puller (model P-97, Sutter Instruments, Novato, CA).

Voltage-gated Ca²⁺ currents (I_{Ca}) were measured using conventional whole-cell voltage-828 829 clamp sampled at a frequency of 50 kHz and low-pass filtered at 2 kHz. Arterial myocytes 830 were continuously perfused with 115 mM NaCl,10 mM TEA-Cl, 0.5 mM MgCl₂, 5.5 mM 831 glucose, 5 mM CsCl, 20 mM CaCl₂, and 10 mM HEPES, adjusted to pH 7.4 with CsOH. 832 Micropipettes were filled with an internal solution containing 20 mM CsCl, 87 mM Aspartic acid, 1 mM MgCl₂, 10 mM HEPES, 5 mM MgATP, and 10 mM EGTA adjusted to pH 7.2 833 834 using CsOH. Current-voltage relationships were obtained by exposing cells to a series of 835 300 ms depolarizing pulses from a holding potential of -70 mV to test potentials ranging 836 from -70 to +60 mV. A voltage error of 9.4 mV due to the liquid junction potential of the recording solutions was corrected offline. Voltage dependence of Ca²⁺ channel activation 837 838 (G/G_{max}) was obtained from the resultant currents by converting them to conductance via 839 the equation $G = I_{Ca}/(\text{test potential} - \text{reversal potential of } I_{Ca})$; normalized G/G_{max} was 840 plotted as a function of test potential. Time constants of activation and inactivation of Ica 841 were fitted with a single exponential function.

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Ikv recordings were performed in the whole-cell configuration with myocytes exposed to
an external solution containing 130 mM NaCl, 5 mM KCl, 3 mM MgCl₂, 10 mM Glucose,
and 10 mM HEPES adjusted to 7.4 using NaOH. The internal pipette solution constituted
of 87 mM K-Aspartate, 20 mM KCl, 1 mM CaCl₂, 1 mM MgCl₂, 5 mM MgATP, 10 mM
EGTA, and 10 mM HEPES adjusted to 7.2 by KOH. A resultant liquid junction potential

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848 of 12.7 mV from these solutions was corrected offline. To obtain current-voltage 849 relationships cells were subjected to a series of 500 ms test pulses increasing from -70 850 to +70 mV. To isolate the different K⁺ channels attributed to composite I_K, cells were first 851 bathed in external I_K solution, subsequently exposed to 100 nM lberiotoxin (Alomone, 852 Jerusalem, Israel) to eliminate any BK_{Ca} channel activity and finally immersed in an 853 external solution containing both 100 nM Iberiotoxin and 100 nM Stromatoxin (Alomone, 854 Jerusalem, Israel) to block both BK_{ca} and K_V2.1 activity. Ionic current was converted to 855 conductance via the equation $G = I(V-E_K)$. E_K was calculated to be -78 mV. Activation 856 time constants for $K_{V2.1}$ currents were obtained by fitting the rising phase of these 857 currents with a single exponential function.

858

859 BK_{ca}-mediated spontaneous transient outward currents (STOCs) and membrane 860 potential were recorded using the perforated whole-cell configuration. To measure both, 861 myocytes were continuously exposed to a bath solution consisting of 130 mM NaCl, 5 862 mM KCl, 2 mM CaCl₂, 1 mM MgCl₂, 10 mM glucose, and 10 mM HEPES, pH adjusted to 863 7.4 with NaOH. Pipettes were filled with an internal solution containing 110 mM Kaspartate, 30 mM KCl, 10 mM NaCl, 1 mM MgCl₂, 0.5 mM EGTA, and 10 mM HEPES 864 865 adjusted to a pH of 7.3 with KOH. The internal solution was supplemented with 250 µg/ml 866 amphotericin B (Sigma, St. Louis, MO). STOCs were measured in the voltage-clamp 867 mode and were analyzed with the threshold detection algorithm in Clampfit 10 (Axon 868 Instruments, Inc). Membrane potential was measured using the current clamp mode.

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870 STOCs were recorded using the perforated whole-cell configuration. The composition of 871 the external bath solution consisted of 134 mM NaCl, 6 mM KCl, 1 mM MgCl₂,2 mM 872 CaCl₂,10 mM glucose, and 10 mM HEPES adjusted to a pH of 7.4 using NaOH. Pipettes 873 were filled with an internal solution of 110 mM K-aspartate, 10 mM NaCL, 30 mM KCl, 1 874 mM MgCl₂, 160 µg/ml amphotericin B, and 10 mM HEPES using NaOH to adjust to pH to 875 7.2. Myocytes were sustained at a holding potential of -70 mV before being exposed to a 876 400 ms ramp protocol from -140 to +60 mV. A voltage error of 12.8 mV resulting from the 877 liquid junction potential was corrected for offline. Kir channels were blocked using 100 µM Ba²⁺. 878

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880 **1.4 Statistics**

Data are expressed as mean ± SEM. All data sets were tested for normality. Normally
 distributed data were analyzed using T-tests or one-way analysis of variance (ANOVA).
 ANOVA analyses were followed by multiple comparison tests (i.e. Tuckey). P<0.05 was
 considered statistically significant.

885

886 Section 2. Computational modeling and simulation

887 2.1 Cell size and structure

The mean capacitance of the cells was experimentally calculated to be 16±3 pF based on all the male and female WT mesenteric C57BL/6J cells utilized in the experiments (N=45). Assuming the cells are roughly cylindrical in shape, the expected radius should be 2.485 μm and a length of 100 μm leading to a surface area of 1.6×10⁻⁵ cm² and a total volume of approximately 1.94×10⁻¹² liters. The cell capacitance of excitable membranes is assumed to be 1.0×10⁻⁶ F/cm², with the calculated surface area the estimated total cell capacitance is C_m = 16 pF.

895

Because the total cell volume is roughly 2×10^{-12} liters, it is assumed that 50% of the total 896 897 cell volume is occupied by organelles. There are three main compartments in vascular 898 myocytes important to the regulation of membrane potential and calcium signaling: the 899 cytosol, sarcoplasmic reticulum (SR), and specialized junctional domains formed by the 900 SR and the plasma membrane. The cytosol occupies approximately 50% of total cell volume ($V_{cvt} = 1.0 \times 10^{-12}$ L). The sarcoplasmic reticulum occupies approximately 5% of 901 902 cell cytosolic volume (V_{SR} = 5.0×10^{-14} L), and the junctional domain volume is approximately 1% of the cytosol volume $(V_{Jun} = 0.5 \times 10^{-14} \text{ L})^{22,37-39}$. 903

904

905 **2.2 Model development**

The male and female *in silico* models are single whole-cell models based on the electrophysiology of isolated mesenteric vascular smooth muscle myocytes. A schematic of the proposed model is shown in **Figure 1**. The membrane electrophysiology can be described by the differential equation:

32

910
911 (22)
$$\frac{dV}{dt} = \frac{-I_{ion}}{C_m}$$

 C_m

912

913 Where V is voltage, t is time, C_m is membrane capacitance I_{ion} is the sum of 914 transmembrane currents. The contribution of each transmembrane current to the total 915 transmembrane ionic current can be described by the following equation:

916

917 (23) $I_{ion} = (I_{KV1.5} + I_{KV2.1} + I_{BKCa} + I_{K,b} + I_{Cav1.2} + I_{PMCA} + I_{Ca,b} + I_{NCX} + I_{NSC} + I_{Na,k} + I_{Na,b})$ 918

919 The eleven transmembrane currents are generated by ion channels, pumps, and 920 transporters. Currents from ion channels include the voltage-gated L-type calcium current 921 (I_{Ca}), the nonselective cation current (I_{NSC}), voltage-gated potassium currents (I_{Kv1.5} and $I_{Kv2,1}$), and the large-conductance Ca²⁺-sensitive potassium current (I_{BKCa}). Additionally, 922 923 there are three background or leak currents ($I_{K,b}$, $I_{Ca,b}$, and $I_{Na,b}$). Currents from pumps 924 and transporters include the sodium-potassium pump current (I_{NaK}), plasma membrane 925 Ca-ATPase transport current (IPMCA), and sodium-calcium exchanger current (INCX).

926

927 Cytosolic concentrations of sodium and potassium as a function of time are determined 928 by considering the sum of their respective fluxes into the cytosol.

929

930 (24)
$$\frac{d[K^+]_{cyt}}{dt} = -\frac{(I_{K\nu2.1} + I_{K\nu1.5} + I_{BKca} + I_{K,b} - 2I_{NaK} + I_{NSC-K})}{z_{K} Vol_{cyt} F}$$

931
932 (25)
$$\frac{d[Na^+]_{cyt}}{dt} = -\frac{(3I_{NCX} + 3I_{NaK} + I_{Na,b} + I_{NSC} - Na)}{z_{Na} Vol_{cyt} F}$$
933

Where F is the Faraday's constant, Vol_{cyt} is the cytoplasmic volume and z_k and z_{Na} are 934 935 the valence of potassium and sodium ions, respectively.

936

937 The calcium dynamics is compartmentalized into three distinct regions: cytosol $[Ca^{2+}]_{i}$. the sarcoplasmic reticulum $[Ca^{2+}]_{SR}$, and the junctional region $[Ca^{2+}]_{jun}$. The cytosol 938 939 includes a calcium buffer, which we assume can be described as a first-order dynamics 940 process.

941

942 *Cytosolic calcium region (*[Ca²⁺]_i): calcium concentration in this region varies between
943 100-300 nM¹³ and is mainly influenced by the following fluxes: transmembrane pumps
944 and transporters, the sarcoplasmic reticulum Ca-ATPase (J_{SERCA}), diffusion from the
945 junctional domain region (J_{Jun-Cyt}) and the calcium buffer calmodulin (BUF_{CAM}).

946

947 (26)
$$\frac{d[Ca^{2+}]_i}{dt} = -\frac{(I_{NCX} + I_{Cav1.2} + I_{Ca,b} - 2I_{PMCA})}{z_{Ca}FV_{Cyt}} - J_{SERCA} + J_{jun-cyt} - \left(k_{BUF_{on}} * Ca_i * Ca_i + J_{inc} + Ca_i + Ca$$

948
$$(BUF_T - BUF_{CAM}) - k_{BUF_{off}} * BUF_{CAM})$$

949

Sarcoplasmic reticulum region ($[Ca^{2+}]_{SR}$): calcium concentration in this region varies between 100-150 μ M⁴⁰ and it is mainly influenced by the sarcoplasmic reticulum Ca-ATPase (J_{SERCA}) and the flux from the ryanodine receptors (J_{RyR}).

953

954 (27)
$$\frac{d[Ca]_{SR}}{dt} = \left[\frac{Vol_{Cyt}}{Vol_{SR}}\right] J_{SERCA} - \left[\frac{Vol_{Cyt}}{Vol_{SR}}\right] \left[J_{Ryr}\right]$$

955

Junctional region ([Ca^{2+}]_{jun}): calcium concentration in this region varies between 10-100 μ M^{37,38} and is mainly influenced by the flux from the ryanodine receptors (J_{RyR}), the diffusion from the junctional region to the cytoplasm (J_{Jun-Cyt})

959

960 (28)
$$\frac{d[Ca]_{jun}}{dt} = \left[\frac{Vol_{Cyt}}{Vol_{jun}}\right] J_{Ryr} - \left[\frac{Vol_{Cyt}}{Vol_{jun}}\right] J_{jun-cyt}$$

961

In the model, the flux of J_{SERCA} was adapted from the Luo-Rudy II model⁴¹ and the flux of J_{RyR} was adapted from previous models of ryanodine receptors activation, originally introduced in the field of cardiac electrophysiology^{42–44}.

965

966 **2.3 Parameter optimization and reformulation of the gating ion channel models**

The ionic current models of I_{Ca}, I_{Kv2.1}, and I_{Kv1.5} were optimized using the approach employed by Kernik *et al.*⁴⁵. Here, the open probability P_o of each voltage-dependent gating variable "n" was defined by opening- and closing-rate voltage-dependent functions α_n and β_n respectively, and were modeled by simple exponentials of the form:

34

971 (29)
$$\alpha_n(V) = x_1 e^{\left(\frac{V}{x_2}\right)}$$

972 (30)
$$\beta_n(V) = x_3 e^{\left(\frac{V}{x_4}\right)}$$

973 (31)
$$\tau_n(V) = \frac{1}{\alpha_n(V) + \beta_n(V)} + x_5$$

974

975 The steady-state availability remains the same as the classical Hodgkin-Huxley 976 formulations and the time constant values follow a modified version formulation by 977 accommodating an extra parameter x_5 in equation 31. $(x_1, x_2, x_3, x_4, x_5)$ are parameters 978 to be optimized using experimental data. We used the parameter optimization employed 979 by Kernik et al.⁴⁵, which minimizes the error between model and experimental data using 980 the Nelder–Mead minimization of the error function. Random small perturbations (<10%) 981 were applied to find local minima, to improve data fit. The parameter fit with the minimal 982 error function value after 1000 to 10000 perturbations was used as the optimal model fit 983 to the data.

984

985 **2.4 Cellular simulations with noise**

The simulations encapsulate the cumulative effect of stochastic ion channel activity on cell voltage dynamics through the fluctuating current term, $\xi(t)$, into the membrane potential (dV/dt) equation⁴⁶, as shown in equation 32. Here it is assumed $\xi(t)$ is only a function of time and it is implemented as Gaussian white noise⁴⁷.

990 (32)
$$\frac{dV}{dt} = -\frac{I_{total}(V)}{C_m} + \sigma \xi(t)$$

We use the Euler-Maruyama numerical method for updating equation 33 and 34 asfollows.

993 (33)
$$V(t + \Delta t) = V(t) - \frac{I(V(t))}{c_m} \Delta t + \sigma * \text{randN} * \sqrt{\Delta t}$$

994 (34)
$$\frac{d[Ca]_{SR}}{dt} = \left[\frac{Vol_{Cyt}}{Vol_{SR}}\right] J_{SERCA} - \left[\frac{Vol_{Cyt}}{Vol_{SR}}\right] [J_{Ryr}] + \sigma * \text{randN} * \sqrt{\Delta t}$$

995 996

Where randN is a random number from a normal distribution (N(0,1)) with mean 0 and variance 1. Δt is the time step and σ is the "diffusion coefficient", which represents the amplitude of the noise. The numerical method for updating the voltage was forward Euler.

1001 **2.5 One-dimensional simulations**

The idealized one-dimensional representation of a vessel was developed by connecting 400 Hernandez-Hernandez model cells in series via simulated resistances to represent gap junctions. For each cell in the cable, the Hernandez-Hernandez model was used to compute ionic currents and concentration changes. The temporal transmembrane fluxes of the Hernandez-Hernandez model are related to the spatial or current flow by a finite difference approximation of the cable equation^{48–50}

1008

1009 (35)
$$\left[C_m \left(\frac{V_i^{(t+1)} - V_i^t}{\Delta t} \right) + I_{ion} + I_{stim} \right] = \frac{a}{4 \left(R_{myo} + \frac{R_g}{\Delta x} \right)} \frac{(V_{i-1}^t - 2V_{i-1}^t + V_{i-1}^t)}{\Delta x \Delta x}$$

where I_{ion} represents the individual membrane ionic current densities (pA/pF) of the Hernandez-Hernandez model, I_{stim} is the stimulus current density (pA/pF) set to zero in our simulation, *a* is the radius of the fiber (5 µm), C_m is the membrane capacity (pA/pF), Vit is membrane potential at segment i and time t, Δx is the discretization element (100 µm = 0.01 cm). Where R_{myo} is the myoplasmic resistance (R_{myo}=150 Ωcm) and R_g is the gap junction resistance (R_g=71.4 Ωcm²).

1016

1017 **2.6 Sensitivity Analysis**

The baseline models in male and female vascular smooth muscle cells were analyzed 1018 1019 through a parameter sensitivity assessment using multivariable linear regression, 1020 following the methodology introduced by Sobie⁵¹. The scope of the sensitivity analysis encompassed variations in the maximal conductance and maximal ion transport rates of 1021 1022 the transmembrane currents, including I_{KV1.5}, I_{KV2.1}, I_{BKCa}, I_{K.b}, I_{Cav1.2}, I_{PMCA}, I_{Ca.b}, I_{NCX}, 1023 INSC, INak, and INa,b. All other parameters, notably those defining model kinetics, remained 1024 constant at the values established by the foundational model. Scaling factors were 1025 randomly selected from a log-normal distribution characterized by a median value of 1 1026 and a standard deviation of 0.1⁴⁵.

1027

1028 **2.7 Simulation protocols**

1029Code for simulations and analysis was written in C++ and MATLAB 2018a. The single1030vascular smooth muscle code was run on an Apple Mac Pro machine with two 2.7 GHz

1031 12-Core Intel Xeon processors and an HP ProLiant DL585 G7 server with a 2.7 GHz 28-1032 core AMD Opteron processor. Vessel simulations were implemented in C++ and 1033 parallelized using OpenMP. The C++ code was compiled with the Intel ICC compiler, 1034 version 18.0.3. Numerical results were visualized using MATLAB R2018a by The 1035 MathWorks, Inc. All codes and detailed model equations are available on GitHub 1036 (https://github.com/ClancyLabUCD/sex-specific-responses-to-calcium-channel-blockers-1037 in-mesenteric-vascular-smooth-muscle)

37

1038

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1291 **FIGURE LEGENDS**

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1294 Figure 1. A schematic representation of the Hernandez-Hernandez model. The 1295 components of the model include major ion channel currents shown in purple including 1296 the voltage-gated L-type calcium current (I_{Ca}), nonselective cation current (I_{NSC}), voltage-1297 gated potassium currents ($I_{KV1.5}$ and $I_{KV2.1}$), and the large-conductance Ca²⁺-sensitive 1298 potassium current (IBKCa). Currents from pumps and transporters are shown in red 1299 including the sodium/potassium pump current (INaK), sodium/calcium exchanger current 1300 (I_{NCX}), and plasma membrane ATPase current (I_{PMCA}). Leak currents are indicated in 1301 green including the sodium leak current ($I_{Na,b}$), potassium leak current ($I_{K,b}$), and calcium 1302 leak current (I_{Ca,b}). In addition, two currents in the sarcoplasmic reticulum are shown in 1303 orange: the sarcoplasmic reticulum Ca-ATPase current (ISERCA) and ryanodine receptor 1304 current (IRVR). Calcium compartments comprise three discrete regions including cytosol 1305 ([Ca]_i), sarcoplasmic reticulum ([Ca]_{SR}), and the junctional region ([Ca]_{Jun}). Red stars (*) 1306 indicate measured sex-specific differences in ionic currents.

1308 Figure 2. Experimentally measured and modeled L-type calcium currents (I_{Ca}) from 1309 male and female vascular smooth muscle cells. Properties of I_{ca} are derived from 1310 measurements in male and female vascular smooth muscle (VSM) cells isolated from the mouse mesenteric arteries following voltage-clamp steps from -60 to 60 mV in 10 mV 1311 1312 steps from a -80 mV holding potential. Experimental data is shown in black circles for male and blue squares for female. Model fits to experimental data are shown with black 1313 1314 solid lines for male and blue solid lines for female. (A) Male and female time constants of 1315 Ica activation. (B) Male and female time constants of Ica inactivation. (C) Male and female 1316 voltage-dependent steady-state activation and inactivation of Ica. (D) Current-voltage (I-V) relationship of I_{Ca} from male and female vascular smooth muscle myocytes. *P < 0.05, 1317 1318 **P < 0.01, ***P<0.001. Error bars indicated mean ± SEM.

Figure 3. Experimentally measured and modeled potassium currents (Ικντοτ) from
 male and female vascular smooth muscle cells. The properties of Ικν1.5 and Ικν2.1 from
 experimental measurements in male and female vascular smooth muscle cells isolated

from the mouse mesenteric arteries were recorded in response to voltage-clamp from -1323 1324 60 to 40 mV in 10 mV steps (holding potential -80mV). Experimental data is shown as 1325 black circles for male and blue squares for female. Model fits to experimental data are 1326 shown with black solid lines for male and blue solid lines for female. (A) Male and female 1327 voltage-dependent steady-state activation of $I_{Ky2,1}$. (B) Male and female time constants of 1328 $I_{Kv2.1}$ activation. (C) Current-voltage (I-V) relationship of $I_{Kv2.1}$ from male and female 1329 myocytes. (D) Male and female voltage-dependent steady-state activation of $I_{KY1.5}$. (E) 1330 Current-voltage (I-V) relationship of $I_{Kv1.5}$ from male and female myocytes. (F) Male and female total voltage-gated potassium current $I_{KVTOT} = I_{KV1.5} + I_{KV2.1}$. (G) Predicted male 1331 1332 and female time constants of the IKV1.5 activation gate. (H) Table showing sex-specific 1333 differences in conductance and steady-state total potassium current-voltage 1334 dependence. *P < 0.05, **P < 0.01, ***P<0.001, ****P<0.0001. Data points without asterisks are not significant. Error bars indicated mean ± SEM. 1335

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Figure 4. Experimentally measured and modeled large-conductance Ca²⁺-activated 1337 1338 K⁺ currents (I_{BKαβ1}). The model was optimized to data from Bao and Cox (Bao & Cox, 1339 2005). (A) Voltage-dependent activation of $I_{BK\alpha\beta1}$ from experiments performed with three 1340 different [Ca]_{Jun} concentrations (1 µM, 10 µM, 100 µM) shown in green circles is the data 1341 from (Zhuge et al., 2002) (B) Voltage-dependent activation time constants with 1342 [Ca]_{Jun}=0.003 µM and simulations [Ca]_{Jun}=10 µM. (C) Simulated I-V curve at different 1343 peak levels of [Ca]_{Jun} levels. (D) Simulated I-V curve with different BK_{ca} average cluster 1344 sizes (N = 4,6, 8, and 10).

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Figure 5. Membrane potential from experiments and simulations in male and female vascular smooth muscle myocytes. A) Whole-cell membrane potential recordings in male and female myocytes showing spontaneous repeat transient hyperpolarization of the membrane potential. **B**) Simulated whole-cell membrane potential with physiological noise. **C**) Comparison of sensitivity analysis performed around the baseline membrane potential in male and female models using multivariable regression.

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Figure 6. Differential effects of voltage-gated potassium current (I_{KVTOT}) block in 1353 1354 male and female myocytes. A) Simulated time course of male I_{Kv2.1} (top panel, solid 1355 traces) and Ikv1.5 (lower panel, dashed traces) at three different baseline membrane 1356 potentials (-45 mV green, -40 mV blue, and -35mV black). B) Simulated time course of female I_{Kv2.1} (top panel solid traces) and I_{Kv1.5} (lower panel, dashed traces) at three 1357 different baseline membrane potentials (-45 mV light blue, -40 mV purple, and -35mV 1358 red). Current contribution to IKVTOT from Kv1.5 (indicated by asterisks) and Kv2.1 in male 1359 1360 and female myocytes at a baseline membrane potential of -45 mV (C), -40 mV (D), and -1361 35 mV (E).

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Figure 7. Simulated L-type calcium currents (Ica) and calcium influx in male and female vascular smooth muscle cells. A) Male and female whole-cell I_{Ca} membrane potential relationship. (B) Male and female intracellular calcium concentration in the cytosolic compartment at indicated membrane potential. C) Time course of membrane potential in male vascular smooth muscle cells before (black) and after (gray) simulated nifedipine application (*top panel*). Corresponding time course of L-type calcium current

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1369 Ica before (black) and after (gray) simulated nifedipine application (*middle panel*) and 1370 intracellular calcium [Ca²⁺]; concentration before (black) and after (gray) simulated 1371 nifedipine application (lower panel). D) Time course of membrane potential in female 1372 vascular smooth muscle cells before (blue) and after (pink) simulated nifedipine 1373 application (top panel). Corresponding time course of L-type calcium current I_{ca} before 1374 (blue) and after (pink) simulated nifedipine application (*middle panel*) and intracellular 1375 calcium [Ca²⁺]_i concentration before (blue) and after (pink) simulated nifedipine 1376 application (lower panel).

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Figure 8. A one-dimensional tissue model representation of vascular smooth 1379 1380 muscle cells connected in series. A) Uncoupled female vessel simulation showing cell 1381 1, cell 50, and cell 100 at a baseline membrane potential of -35 mV. B) Uncoupled male vessel simulations showing cell 1, cell 50, and cell 100 at a baseline membrane potential 1382 1383 of -45 mV. C) Composite female (blue trace) and male (black trace) membrane potential 1384 of 400 coupled smooth muscle cells connected with gap junctional resistance of 71.4 1385 Ω cm² in a one-dimensional tissue representation. (**D**) Sharp-electrode records of the 1386 membrane potential of smooth muscle in pressurized (80-mmHg) female and male 1387 arteries from O'Dwyer et al., 2020.

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1390 Figure 9. Experimentally measured and modeled intracellular calcium [Ca]_i in male 1391 and female vessels and response to clinically used L-type Ca^{2+} channel blocker. A) 1392 Intracellular calcium [Ca]_i in female (blue symbols) and male (black symbols) arteries at 1393 intravascular pressures ranging from 20 to 120 mmHg. Simulations showing [Ca]_i in the 1394 idealized female and male vessels are shown with blue and black solid lines, respectively. Simulated [Ca]; after the application of clinically used L-type Ca²⁺ channel blocker 1395 1396 nifedipine is shown with dashed lines for male (black) and female (blue). B) Comparison 1397 of the percentage change of [Ca]_i in male (black) and female (blue) after the application L-type Ca²⁺ channel blocker nifedipine at 80 mmHg and 120 mmHg. *P < 0.05. **P < 0.01. 1398 1399 1400

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FIGURES AND LEGENDS



Figure 1. A schematic representation of the Hernandez-Hernandez model. The components of the model include major ion channel currents shown in purple including the voltage-gated L-type calcium current (I_{Ca}), nonselective cation current (I_{Nsc}), voltage-gated potassium currents ($I_{Kv1.5}$ and $I_{Kv2.1}$), and the large-conductance Ca²⁺-sensitive potassium current (I_{BKCa}). Currents from pumps and transporters are shown in red including the sodium/potassium pump current (I_{NaK}), sodium/calcium exchanger current (I_{NcX}), and plasma membrane ATPase current (I_{PMCA}). Leak currents are indicated in green including the sodium leak current ($I_{Na,b}$), potassium leak current ($I_{K,b}$), and calcium leak current ($I_{Ca,b}$). In addition, two currents in the sarcoplasmic reticulum are shown in orange: the sarcoplasmic reticulum Ca-ATPase current (I_{SERCA}) and ryanodine receptor current (I_{RyR}). Calcium compartments comprise three discrete regions including cytosol ([Ca]_i), sarcoplasmic reticulum ([Ca]_{SR}), and the junctional region ([Ca]_{Jun}). Red stars (*) indicate measured sex-specific differences in ionic currents.

Figure 2



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Figure 3. Experimentally measured and modeled potassium currents (IKVTOT) from male and female vascular smooth muscle cells. The properties of I_{Kv1.5} and I_{Kv2.1} from experimental measurements in male and female vascular smooth muscle cells isolated from the mouse mesenteric arteries were recorded in response to voltage-clamp from -60 to 40 mV in 10 mV steps (holding potential -80mV). Experimental data is shown as black circles for male and blue squares for female. Model fits to experimental data are shown with black solid lines for male and blue solid lines for female. (A) Male and female voltage-dependent steady-state activation of $I_{Kv2.1}$. (B) Male and female time constants of $I_{Kv2.1}$ activation. (C) Current-voltage (I-V) relationship of $I_{Kv2.1}$ from male and female myocytes. (D) Male and female voltage-dependent steady-state activation of $I_{Kv1.5}$. (E) Current-voltage (I-V) relationship of $I_{Kv1.5}$ from male and female myocytes. (F) Male and female total voltage-gated potassium current $I_{KVTOT} = I_{KV1.5} + I_{KV2.1}$. (G) Predicted male and female time constants of the IKV1.5 activation gate. (H) Table showing sex-specific differences in conductance and steady-state total potassium current-voltage dependence. *P < 0.05, **P < 0.01, ***P<0.001, ****P<0.0001. Data points without asterisks are not significant. Error bars indicated mean ± SEM.





Figure 4. Experimentally measured and modeled large-conductance Ca²⁺-activated K⁺ currents (I_{BKαβ1}). The model was optimized to data from Bao and Cox (Bao & Cox, 2005). (**A**) Voltage-dependent activation of I_{BKαβ1} from experiments performed with three different [Ca]_{Jun} concentrations (1 µM, 10 µM, 100 µM) shown in green circles is the data from (Zhuge *et al.*, 2002) (**B**) Voltage-dependent activation time constants with [Ca]_{Jun}=0.003 µM and simulations [Ca]_{Jun}=10 µM. (**C**) Simulated I-V curve at different peak levels of [Ca]_{Jun} levels. (**D**) Simulated I-V curve with different BK_{ca} average cluster sizes (N = 4,6, 8, and 10).

Figure 5



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Figure 6



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Figure 9
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