# Dual Stretch Responses of mHCN2 Pacemaker Channels: Accelerated Activation, Accelerated Deactivation

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ABSTRACT Mechanoelectric feedback in heart and smooth muscle is thought to depend on diverse channels that afford myocytes a mechanosensitive cation conductance. Voltage-gated channels (e.g., Kv1) are stretch sensitive, but the only voltagegated channels that are cation permeant, the pacemaker or HCN (hyperpolarization-activated cyclic nucleotide-gated) channels, have not been tested. To assess if HCN channels could contribute to a mechanosensitive cation conductance, we recorded I<sub>HCN</sub> in cell-attached oocyte patches before, during, and after stretch for a range of voltage protocols. I<sub>mHCN2</sub> has voltage-dependent and instantaneous components; only the former was stretch sensitive. Stretch reversibly accelerated hyperpolarization-induced I<sub>mHCN2</sub> activation (likewise for I<sub>spHCN</sub>) and depolarization-induced deactivation. HCN channels (like Kv1 channels) undergo mode-switch transitions that render their activation midpoints voltage history dependent. The result, as seen from sawtooth clamp, is a pronounced hysteresis. During sawtooth clamp, stretch increased current magnitudes and altered the hysteresis pattern consistent with stretch-accelerated activation and deactivation. I<sub>mHCN2</sub> responses to step protocols indicated that at least two transitions were mechanosensitive: an unspecified rate-limiting transition along the hyperpolarization-driven path, mode I<sub>closed</sub> → mode II<sub>open</sub>, and depolarization-induced deactivation (from mode I<sub>open</sub> and/or from mode II<sub>open</sub>). How might this affect cardiac rhythmicity? Since hysteresis patterns and "on" and "off" I<sub>HCN</sub> responses all changed with stretch, predictions are difficult. For an empirical overview, we therefore clamped patches to cyclic action potential waveforms. During the diastolic potential of sinoatrial node cell and Purkinje fiber waveforms, net stretch effects were frequency dependent. Stretch-inhibited (SI) I<sub>mHCN2</sub> dominated at low frequencies and stretch-augmented (SA) I<sub>mHCN2</sub> was progressively more important as frequency increased. HCN channels might therefore contribute to either SI or SA cation conductances that in turn contribute to stretch arrhythmias and other mechanoelectric feedback phenomena.

### INTRODUCTION

Mechanical forces affect the electrophysiology of cardiac and smooth muscle cells. Prime candidates for force transduction in mechanoelectric feedback are channels whose open probability changes reversibly with membrane stretch, i.e., mechanosensitive (MS) channels. MS channels include (along with TRP, 2PDK, and others) voltage-gated channels (1). Here, we examine stretch responses of hyperpolarization-activated cyclic nucleotide-gated (HCN) channels, the only cation-selective voltage-gated channels (2). HCN tetramers form pacemaker channels and the subunits have a Kv-like arrangement (six transmembrane-spanning domains each with S1–S4 sensor modules and S5–S6 pore modules).

Highlighting the need to test pacemaker channel mechanosensitivity are the following observations: 1), cell swelling reversibly increases  $I_{mHCN2}$  (3); 2), swelling of ventricular myocytes, cells now known to express HCN channels (e.g., (4,5)), activates an unidentified inwardly rectifying  $I_{cation}$  (6); 3), hypothalamic osmosensory neurons have a swelling- and voltage-dependent Cs-blockable  $I_{cation}$  (7); 4), cholesterol is a bilayer mechanical reagent that modulates voltage-gated channels (8) and altered cholesterol levels in sinoatrial node

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(SAN) cells alter the kinetics of HCN4-based pacemaker channels (9); 5), inhibitors of another type of MS cation conductance that is a candidate mechanotransducer for mechanoelectric feedback, the MS TRP channels, inhibit neither mouse SAN cell MS  $I_{\text{cation}}$  (10) nor certain ventricular mechanoelectric feedback phenomena (11); 6), repeated attempts to record MS TRP-like events from adult cardiomyocytes have failed (12); and 7), many MS cells and MS cellular structures are abundantly endowed with HCN channels (Table 1).

Information on native cell MS voltage-gated channel responses is sparse, but in smooth muscle and SAN cells, voltage-gated calcium channel current reversibly increases with stretch (1) (in recombinant calcium channels this trait resides with the pore subunit (13)). MS modulation of voltagegated channels is best understood for Kv channels, where voltage sensing, slow inactivation, and a concerted step before pore opening have all been characterized as MS transitions (14,15). Reversible stretch responses in these Kv channels occur in both excised and cell-attached patches (16); and since patch excision destroys membrane skeleton organization, we infer that the bilayer transmits force. Why does Ky gating respond to bilayer stretch? We hypothesize that each conformation of a Kv channel's extensive protein/ bilayer interface (17) has a sufficiently different lateral pressure profile (18) that bilayer mechanical perturbations change the relative stabilities of the conformations. Lateral pressure profiles can be perturbed physically (i.e., bilayer expansion/

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#### TABLE 1 HCN-based channels in mechanosensory/ motile structures

Osmosensory neurons-hypothalamic osmosensory magnocellular neurons
of the supraoptic nucleus have a pharmacologically identified I <sub>h</sub>
(pacemaker current) (49); although osmosensitivity requires expression
of a truncated vanilloid receptor (43), the identity of the SI osmosensitive
background cation conductance is unresolved
Demonscripton terminals (50) UCN1 2 and 4 isoforms are averaged in

- Baroreceptor terminals (50)—HCN1, 2, and 4 isoforms are expressed in mechanoreceptor endings of aortic arch mechanosensory neurons
- Sperm flagella (21)—immunocytochemistry (sea urchin isoforms) locates spHCN expression in the sperm flagellum and not in the head. Human testis is HCN4-rich, but cellular, and the subcellular localization in that tissue is unknown (51)
- Vertebrate saccular hair cells (52)—in situ hybridization (subcellular localization unknown)
- Primary afferents/pain (re: tactile allodynia, ectopic discharge) (53) pharmacological inhibition of pacemaker current relieves tactile allodynia. Centrally, HCN2 widely colocalizes with substance P (54).
- Urinary bladder afferents (L6-S1 spinal chord, dorsal root ganglion neurons) (HCN2 is the predominant type of HCN channel expressed;

possible involvement in the micturation reflex) (55)

Primary sensory afferents/enteric nervous system (56)

 $I_{\rm h}$  currents and positive immunoreactivity for HCN1, 2, and 4 but not for HCN3

Dorsal root ganglion sensory afferents

Cell bodies have I<sub>h</sub>, are immunopositive for HCN1 and HCN2 (57)

compression) and/or chemically (e.g., depletion/enrichment of surface-active lipids, alcohols, anesthetics, and drugs). Since the "bilayer mechanics" of membrane stretch and the bilayer mechanics of cholesterol, lysophospholipids, short chain alcohols, phenothiazines, etc. are related forms of lipid stress (19), the fact that HCN channels can be modulated by stretch has wide potential physiological and pathophysiological implications.

Our principal subject here, mHCN2, is known to participate in SAN cell and neuronal rhythmicity through the phenotypes of a knockout mouse (20). We also briefly tested spHCN (21) (from sea urchin) since this is the only isoform for which gating current information has been obtained (22) and the one for which it was shown that HCN and Kv1 voltage sensors respond similarly to voltage (23). In HCN channels, however, it is hyperpolarization, not depolarization, that triggers channel opening. In Kv1 channels, depolarization and stretch facilitate the same step, namely, the independent depolarization-driven activation transition (15). On the face of it, if HCN channels respond like Kv1 channels, then stretch should slow  $I_{\text{HCN}}$  activation and speed  $I_{\rm HCN}$  deactivation. However, that is not the full story. Kv1 channel mutants with identified rate-limiting transitions reveal additional MS transitions: stretch accelerates slow inactivation (15) and it decelerates a concerted voltage-dependent step just before pore opening (14). If comparable transitions occur in HCN channels (not yet established in the case of concerted transitions), they may be similarly MS.

Mode switching, a kinetic trait whose consequences become striking during prolonged voltage excursions, is a feature common to HCN and Kv channels (22). The Kv1 version is termed "C-type inactivation". Depending on recent voltage history, Kv1 channels are in "active" or "slowinactivated" mode; "active" has open and closed states, "slow-inactivated", only closed states. In HCN channels, by contrast, both modes have open and closed states and hence discrete g (V) curves (Fig. 1 A). Mannikko et al. hypothesize (22) that once a HCN channel opens, a voltage-independent conformation change occurs at the pore/sensor domain interface that stabilizes the voltage sensor in its retracted position and thus gives rise to a depolarizing shift in the Q (V) and activation curves. Mammalian HCN1, 2, and 4 isoforms all exhibit mode-switch behavior which, for all of them, occurs on a 100-ms timescale even though activation kinetics vary by well over an order of magnitude (HCN1 is fastest, HCN4 is slowest) (24). The resulting hysteresis in the HCN channel current/voltage (I/V) relations is most prominent in HCN1 and least in HCN4 (25). Prolonged depolarization favors (stabilizes) mode I and prolonged hyperpolarization favors mode II (Fig. 1 A) with the overall consequence that open HCN channels experience longer openings and closed ones, longer closings.

Cardiomyocytes express mixtures of isoforms HCN1, 2, and 4 (26). Smooth muscle shows electrophysiological and reverse transcription-polymerase chain reaction evidence of HCN2, 3, and 4 (27). In mammalian brain, all four isoforms are expressed, each with a unique pattern of distribution (28); immunogold electronmicroscopy shows that, as well as being expressed in neurons, HCN2 has a widespread expression in oligodendrocytes (29). Native HCN-based channels, with conductances of <10 pS, pass Ca<sup>2+</sup> (30) as well as Na<sup>+</sup> and K<sup>+</sup>.

We expressed HCN channels in oocytes, monitoring macroscopic  $I_{\rm HCN}$  in cell-attached patches before, during, and after stretch, using a variety of voltage clamp protocols. In Kv1 channels depolarization elicits activation of closed channels, and the process accelerates with stretch (15). In HCN channels, depolarization elicits deactivation of open channels, and we predicted it too would accelerate with stretch since it involves comparable motions of the voltage sensors (23). Thinking of HCN channels as "inverted Kv1 channels" we therefore expected to observe "stretch inhibition" (SI) during hyperpolarization (due to stretch-enhanced deactivation). We were therefore surprised to observe a very unequivocal "stretch augmentation" (SA). Eventually we found that, indeed, deactivation was accelerated with stretch, but this strong effect was initially cryptic, since a), it did not result in a slower activation of  $I_{\rm HCN}$  during hyperpolarizations, and b), we did not immediately understand the workings of HCN channels protracted tail currents.

Having unexpectedly found that HCN current activation accelerated with stretch, a second surprise came when we tested currents during rhythmic activity. During slow action potential (AP) waveforms, the SI effects, not the SA effects, were what dominated. More particularly, during the diastolic part of APs,  $I_{mHCN2}$  showed SI at low and SA at high "beat"



FIGURE 1 Modes and a common kinetic framework for HCN and Kv channels. (A) After prolonged sojourns ( $\infty$ ) at hyperpolarized or depolarized voltages, HCN channels principally occupy mode II (gray) or mode I (black) (22), respectively. Activation and gating charge curves (g (V), Q (V)) for the modes are depicted. The vertical arrow depicts the consequence (at a given voltage) of undergoing a mode I to mode II transition. (B) Open (O) and closed (C) states connected in circular schemes (22) for HCN and Kv channels (d, depolarization; h, hyperpolarization). The x through the O<sub>I</sub> of Kv signifies that in Kv-WT, "OI", unlike its HCN counterpart, is occluded (i.e., "inactivated-nonconducting") (see Discussion). The dominant states for typical V<sub>hold</sub> potentials (depolarized-HCN; hyperpolarized-Kv) are boxed. Asterisks and inverse asterisks signify candidate stretch-accelerated and stretch-decelerated transitions, respectively, that would represent a consistent pattern across subfamilies. Two additional candidates for stretchaccelerated transitions bear question marks (see Discussion). The stretchdecelerated (1/\*) Kv transition (14) is the concerted depolarization-driven closed-closed transition just before the last (voltage-independent) pore opening step (67,68). Whether HCN channels have analogous concerted transitions (shown gray) into the open state(s) is unknown. The asterisks in mode II correspond to a known stretch-accelerated voltage sensor motion for Kv (15); evidence suggesting that the comparable mode I transition is also stretch-accelerated is presented here for mHCN2.

frequencies. It required the emerging understanding of HCN channel mode switch and the hysteresis that results from mode switch (22) to help clarify the time-dependent stretch responses of the mHCN2 pacemaker channels.

### **METHODS**

#### Oocyte preparation and cRNA injection

*Xenopus laevis* oocytes were injected with 5–55 ng cRNA as described previously (14). Capped cRNA for mHCN2 (mHCN2-pGEM-HE (kindly provided by Steve Siegelbaum); linearized with SphI) (31) and spHCN (21) (SPIH-pGEM-HE (kindly provided by U. B. Kaupp (21)); linearized with

NheI) was produced by in vitro transcription using Ambion message machine.

### Electrophysiology

Manually devitellinated oocytes were transferred to an inverted microscope rig. Thick-walled, sylgard-coated, fire-polished (using a soda glass-coated platinum filament) pipettes were prepared as previously (14) and had resistances of 2–4 M $\Omega$ . Macroscopic currents measured from cell-attached patches (Axopatch 200B; Axon Instruments, Foster City, CA) at RT (20°C–22°C) were filtered at 5 kHz (Axopatch 200B low-pass filter) and analyzed using WinASCD (Guy Droogmans) and Origin (Microcal, Northampton, MA).

Experimental voltage protocols (step, ramp, and waveform protocols) were controlled via pClamp 8, as indicated in Results.  $V_{hold}$  was 0 mV unless otherwise noted.

Membranes were stretched reversibly via negative pipette pressure, controlled, and monitored by a DPM-1B transducer (Bio-Tek Instruments, Winooski, VT) or a high-speed pressure clamp (HSPC-1, ALA Scientific Instruments, Westbury, NY). MS responses were typically elicited using -30 or -40 mmHg. Because a given pressure generates a different tension in different patches, several levels were sometimes tested. To obtain difference currents, protocols were structured as Before/During/After stretch (*B/D/A*) sets, with at least 15 s between runs within a *B/D/A* set. Data were discarded when stretch-induced changes were irreversible.

Water-injected (in actuality, Tris buffer-injected) control oocytes were incubated for 2–3 days, patched and clamped using protocols like those used for Figs. 2–88 (i.e., step, ramp, and AP clamp protocols) to make B/D/A sets (using –40 mm Hg or more). The same lanthanum (1 mM), high-K pipette solution was used for control oocytes as for HCN expressing oocytes. The control records (available online as Supplementary Material) showed virtually no evidence of time- and voltage-dependent conductances.

### Solutions

High-K solution (for bath and pipette) contained (in mM) 89 KCl, 0.4 CaCl<sub>2</sub>, 5 HEPES, 0.8 MgCl<sub>2</sub> (pH 7.5 with KOH). Then 40  $\mu$ M GdCl<sub>3</sub> or 1 mM LaCl<sub>3</sub> was added to pipettes to block the endogenous MS cation channels. Lanthanum (adopted partway through the project) was always fully effective. Gadolinium was sometimes fully effective, but sometimes MS cation channel activity made records unusable, despite adding degassed acidified stock solution (100 mM GdCl<sub>3</sub>) to degassed pipette solution immediately before recording (32). Either gadolinium precipitated imperceptibly or oocytes sporadically express La-sensitive-Gd-insensitive  $I_{MScation}$  channels; renal cells (33) reportedly have such channels.

### RESULTS

# Monitoring recombinant HCN channels against an *I*<sub>MScation</sub> background

To illustrate why lanthanides were needed, Fig. 2 Ai shows a "worst-case" example of interference by endogenous  $I_{\rm MScation}$ . A hyperpolarizing ramp was applied before, during, and after stretch (B/D/A). Here and throughout, black/red/ gray signify B/D/A. The B and A traces run along the x axis, and  $I_{\rm MScation}$  (i.e., the current elicited by stretch, the D trace) corresponds to ~40 open MS<sub>cation</sub> channels. In Fig. 2 Aii, as in 2 Ai, -100-mmHg suction was used, but there was no stretch current because 40  $\mu$ M gadolinium in the pipette inhibited  $I_{\rm MScation}$ .  $I_{\rm HCN}$  families in the presence of lanthanides are shown in Fig. 2 Bi and ii (mHCN2; 40  $\mu$ M



FIGURE 2 Dealing with the endogenous MS cation channel. (A) Ramp (33 mV/s) currents on patches from uninjected oocytes before, during, and after (B/D/A: black, red, gray) stretch without (i) or with (ii) 40  $\mu$ M Gd<sup>3+</sup> in the pipette. Typically, stretch was applied using less intense suction than the -100mmHg used here. (B) spHCN1 (i) and ImHCN2 (ii and iii) current families, using 20-mV steps from a V<sub>hold</sub> of 0 mV (i) or +20 mV (*ii*, *iii*) to -140 mV (*i* and *iii*) or -160 mV (*ii*), with 40  $\mu$ M Gd<sup>3+</sup> (*i*, ii) or 1 mm La<sup>3+</sup> (iii) in the pipette; scales 200 pA/0.5 s; 100 pA/0.25 s; and 200 pA/0.2 s (i-iii). (C) A typical inwardly rectifying ramp I<sub>HCN</sub> (33 mV/s) for mHCN2; arrow indicates ramp direction. (D) Ramp currents (as in C) with a B/D/A protocol, using strong suction (-80 mmHg); in the  $I_{\text{diff}} = D - [(B + 1)^{-1}]$ A)/2], the small instantaneous component gets removed.

gadolinium and 1 mM lanthanum chloride, respectively) and 2 *Biii* (spHCN; 40  $\mu$ M gadolinium chloride). A hyperpolarizing ramp (Fig. 2 *C*) reveals the minor ohmic component reversing at ~0 mV (comprising the instantaneous component (34)  $I_{\text{HCNinst}}$ , nonspecific  $I_{\text{leak}}$ , and any imperfectly inhibited  $I_{\text{MScation}}$ ) and  $I_{\text{HCN}(V,t)}$  for mHCN2 channels, most of which would have been in mode I as they activated at ~-100 mV. The *B/D/A* ramp traces in Fig. 2 *D* show that inhibiting  $I_{\text{MScation}}$  (as in Fig. 2 *Bii*) allowed for extraction of a stretch difference current,  $I_{\text{diff}}$  (i.e., D - [(B + A)/2]), shown in pink.

Capacitive transients interfered with early  $I_{\text{HCN}}$  during voltage steps (e.g., Figs. 3 *A* and 4 *A*). Nevertheless, we used no online linear subtraction since that requires long excursions to mode-switch-inducing voltages and would obliterate any stretch changes in  $I_{\text{HCNinst}}$ . Traces are either raw current, averaged replicates (2 or 3), or  $I_{\text{diff}}$  (i.e., D - [(B + A)/2] from a B/D/A protocol). "Stretch-augmented" (SA) and "stretch-inhibited" (SI) correspond to "inward" and "outward"  $I_{\text{diff}}$ , respectively.

A priori, stretch might alter unitary HCN channel conductance. Since smooth muscle (35) cells show large (>50 pS) hyperpolarization- and stretch-activated  $I_{\text{cation}}$  events, sporadic unitary events riding atop macroscopic  $I_{\text{mHCN2}}$  during stretch (e.g., Fig. 3 *B*) were of interest. However, brief depolarizations showed that such events never inwardly rectified, and since they never occurred with La<sup>3+</sup> (see Methods), we take them to be endogenous currents.

### Stretch increases *I*<sub>HCN</sub> during hyperpolarizing steps

Fig. 3 *A* shows that hyperpolarization-induced  $I_{mHCN2}$  was accelerated and augmented by stretch. As is common for recombinant  $I_{HCN}$ , sporadic inward noise (e.g., as in *red* and

gray traces) was sometimes seen at large hyperpolarizations but was unrelated to stretch. At  $V_{hold} = -40$ mV there was no  $I_{HCN}$  (note I = 0 pA at the start of traces). After prolonged hyperpolarizations, however, stepping back to -40 mV produced a sustained tail current. In other words, the 80-mV drop produced, as expected, an ohmic decrease in  $I_{HCN}$ , but this was not followed by decay toward I = 0. Channels stayed open. We return later to this manifestation that mode switching had occurred during the hyperpolarized sojourn.

As seen in Fig. 3 *B*, even with Gd<sup>3+</sup> in the pipette, patches sometimes exhibited inward and outward unitary  $I_{\rm MScation}$ events. The events contaminating two of the stretch traces illustrate a stark difference between MS  $I_{\rm HCN}$  and endogenous  $I_{\rm MScation}$ : even at the few pA level, MS  $I_{\rm HCN}$  was "macroscopic". Moreover, endogenous unitary  $I_{\rm MScation}$ was not time dependent (note the unitary events during the latent period for the voltage- and time-dependent component of  $I_{\rm mHCN2}$ ) (see also Fig. 5 *A* (*x*, bottom)).

# Stretch does not affect the instantaneous component of $I_{mHCN2}$

Not only did stretch not induce inwardly rectifying high conductance states of the mHCN2 channels, it did not alter  $I_{mHCN2inst}$ . This was directly evident from the insensitivity to stretch of  $I_{mHCN2inst}$  in records where the voltage- and time-dependent component of  $I_{mHCN2}$  increased with stretch. This lack of effect is apparent by inspection of the latent periods for  $I_{mHCN2}$  in Fig. 3 *B* (see also Fig. 5 *C* and Fig. 6, *A* and *B*) and by inspection of hyperpolarizing ramps without and with stretch in Fig. 2 *D* (see also Fig. 7, *B* and *C*).

### Activation of spHCN is stretch sensitive

In B/D/A sets,  $I_{spHCN}$  (Fig. 4 A) exhibited stretch-accelerated activation (*arrows*). As a semiquantitative measure of the



FIGURE 3 Stretch, oocyte patches, mHCN2, and the subcomponents of  $I_{\rm mHCN2}$  40  $\mu$ M Gd<sup>3+</sup> present.  $V_{\rm hold} = -40$  mV. (A) Averaged  $I_{\rm mHCN2}$  for two sequential B/D/A sets (D, -30 mmHg). Notable are the large instantaneous current component (evident at -120 mV above the *dashed line*) and the effect of stepping back to  $V_{\rm hold}$ .  $I_{\rm diff}$  for the averaged sets is plotted at right. The asterisk is mentioned in the text. (B) Five sequential B/D/A sets (D, as indicated in mmHg). The voltage protocol was to +40 mV then to -120 mV as in A. The unitary currents are discussed in the text.

effect of stretch (using -30 mmHg) note that, for more than 150 ms, more than half of the time- and voltage-dependent component of  $I_{spHCN}$  comprised SA  $I_{HCN}$  (Fig. 4 *B*).  $I_{diff}$  peaked at ~150 ms for all three stretch intensities tested (Fig. 4 *C*). Qualitatively, the traces show unequivocal reversible acceleration of activation by stretch, but as the expanded  $I_{diff}$  sections show, the early difference currents were too nonuniform to allow for standardized curve fitting and assignment of rates (as was true, too, for mHCN2 activation currents).

# Stretch-augmented and stretch-inhibited $I_{mHCN2}$ : current activation and tail currents

Fig. 5 *A* shows  $I_{mHCN2}$  for an unusually stable patch: first a control set, then three *B/D/A* sets (with D = -20 mmHg; these are averaged in Fig. 5 *B*). (We did not attempt to study tail currents at positive depolarized voltages; the brief steps

to +40 mV were internal controls to ensure that endogenous MS channels were inhibited.) Stretch-accelerated activation resulted in an elevated steady-state  $I_{mHCN2}$ . On stepping to -40 mV, the magnitude of tail  $I_{diff}$  (*pink arrow*) reflects the stretch augmentation of current that had occurred at -110 mV. The persistence of  $I_{diff}$  at -40 mV (which, note was also  $V_{hold}$ ) suggests that mode switch had occurred during the sojourn at -110 mV (at hyperpolarized voltages, the channels prefer mode II with its right-shifted g (V)). Stretch evidently reversibly affected a rate-limiting transition along the pathway mode  $I_{closed} \rightarrow$  mode II<sub>open</sub> (i.e., the path that is suggested by the vertical arrow in Fig. 1 *A*). Longer tail currents than obtained here were needed to clarify the shape and polarities of  $I_{diff}$  (Fig. 5 *B*\*).

At -40 mV, tail currents required ~2 s for completion, as seen in Fig. 6 A. At right, the  $I_{\text{diff}}$  for the D = -50 mmHg B/D/A set is shown below that for the D = -30 mmHg set (acquired before the -50 mmHg set). Tail currents exhibited an open-channel latency (~100 ms) after which stretch reversibly accelerated the decay of  $I_{\text{HCN}}$  to zero. Note that tail  $I_{\text{diff}}$  (*pink*) was initially stretch augmented, but then crossed the zero current line and became stretch inhibited (likewise, Fig. 6 C\*). The inflection (after ~0.5 s) in the Fig. 6 A tail  $I_{\text{diff}}$  simply indicates the approach to steady state (likewise for spHCN, Fig. 4, for inflections in the rising phase  $I_{\text{diff}}$  and for the rising phase  $I_{\text{diff}}$  here at -140 mV).

Stretch acceleration of  $I_{mHCN2}$  deactivation was robust. In the four B/D/A sets of Fig. 6, B-D, the hyperpolarizing prepulse was successively deepened (-80, -100, -120,-140 mV). At the onset of the -40 mV tail currents, the post -140 mV case (Fig. 6 D, bottom) should have the greatest fraction of channels in mode II<sub>open</sub>. Whether or not the latent period, which is clearly evident as the flat portion in the Fig. 6 C tail  $I_{\text{diff}}$ , was stretch sensitive is uncertain. The shorter latency here at -50 mm Hg than at -30 mmHg in Fig. 6 A (see the  $I_{\text{diff}}$  traces) suggests that it might be. The latency presumably included "electrically silent" mode  $II_{open} \rightarrow$ mode I<sub>open</sub> transitions and (as per the expanded version of the Mannikko et al. model (22)) independent S4 motions between the (multiple) open states of either mode. The deactivation phase, which would include mode  $I_{open} \rightarrow mode$  $I_{closed}$  and mode  $II_{open} \rightarrow mode \ II_{closed}$  transitions, was, however, unequivocally accelerated by stretch. This is the response depicted by the asterisk in Fig. 1  $B^*$ , HCN.

Thus, stretch accelerated  $I_{mHCN2}$  activation during hyperpolarization and  $I_{mHCN2}$  deactivation during depolarization. In mHCN2, therefore, at least two types of transition (with opposing effects on current) were susceptible to the bilayer mechanical perturbation (36) produced by membrane stretch.

Our data showed that the "sign" (SA versus SI) of  $I_{\text{diff}}$  at a given voltage (and time) depended on voltage history. We emphasize this because it would be crucial in designing B/D/A stretch tests for any native  $I_{\text{pacemaker}}$  and in interpreting the outcomes. Consider Fig. 6 *D*, where a 0.7-s depolarization to -40 mV had four different "histories". During that 0.7 s,



FIGURE 4 spHCN1 and stretch. (*A*). Three *B/D/A* sets recorded (same patch) in the sequence shown. The protocol was a simple step from 0 mV to -80 mV. The consistently smaller amplitude of the after traces (*gray*) within each *B/D/A* and the progressive decrease in the amplitudes of before traces (*black*) from one *B/D/A* indicates that slow rundown was occurring, but this did not prevent stretch acceleration of activation. (*B*) Expanded early part of the first *B/D/A* set (i.e., D = -30 mmHg; *black trace* is (B + A)/2) with vertical red and black bars of equal amplitude. (*C*) *I*<sub>diff</sub> for these *B/D/A* sets, with early section expanded. Scale bars (*top* to *bottom*); 100 pA/150 ms/, 30 pA/400 ms, 30 pA/60 ms.

 $I_{\rm diff}$  was mostly SA post -80 mV but mostly SI post -140mV. With mode switching plus stretch acceleration (on both the activation and deactivation pathways) taken into account, this makes sense. Where there is net SA during hyperpolarization, the Idiff must start out SA, but given time they all become SI current (post -80 mV, 0.7 s was clearly not sufficient time). Except for being prematurely truncated, the Fig. 6 D tail currents behave as in Fig. 6 A (likewise for Fig. 3  $A^*$  and Fig. 5\*). Consider, in Fig. 6 A, the small SI  $I_{\text{diff}}$  at -80 mV versus, in Fig. 6 B, the (eventual) SA  $I_{\text{diff}}$  at -80mV. This too is perfectly understandable in terms of mode switch plus dual stretch effects but would be perfectly confusing if voltage history was ignored. Specifically, after a sojourn (>1 s) at -40 mV, mode  $I_{\text{closed}}$  would dominate, but (as the SI traces for  $I_{\text{diff}}$  at -30 mmHg and -50 mmHg Fig. 6 A reveal) stretch-accelerated deactivation caused further closure. Consequently, during stretch, a step to -80 mV produced what was initially an SI Idiff. Given sufficient time at -80 mV, as seen in Fig. 6 B, activation (presumably into mode II<sub>open</sub> far more than into mode I<sub>open</sub>) began and stretchaccelerated activation began to overwhelm the effect of stretch on deactivation, thus producing the net SA effect.

### Voltage hysteresis: dual effects of stretch during sawtooth ramps

HCN voltage hysteresis (25) during sawtooth ramp clamp is typified by right-shifted, more S-shaped I/V relations in depolarizing limbs (22). For HCN channels, unlike Kv channels, moderate slowing of ramp speeds fails to make hyperpolarizing and depolarizing limbs converge (22). Though hysteresis is less pronounced in mHCN2 than in mHCN1 (24,25), mHCN2 (with 1 mM La in the pipette) nevertheless showed the characteristic pattern noted for mHCN1 (i.e., increased rather than decreased hysteresis as ramps are made moderately slower (22)), as seen in Fig. 7 A. The hysteresis persisted during stretch (Fig. 7 B). For the first (hyperpolarizing) limb, applying stretch was essentially like running a slower ramp;  $I_{mHCN2}$  increased. For the (second) depolarizing limb, stretch and ramp speed (= voltage history) interacted strongly (Fig. 7 Ci and ii). At first glance, only SA current is apparent, but that reflects the voltage history of the ramp protocol: the sawtooth ramps, like typical HCN step protocols, started from depolarized voltages. Fig. 7 Ciii shows a normalized plot of stretch and no-stretch traces for the slower (i.e., closer to equilibrium) sawtooth pair. This plot demonstrates, in relative terms, clear evidence of stretch-accelerated deactivation of  $I_{mHCN2}$  (i.e., clear evidence of "SI behavior") in the depolarizing limb. In some patches (not shown) the effect of stretch on deactivation was sufficient to yield outward difference current in the depolarizing limb. Thus, step and sawtooth protocols yielded a consistent pattern of SA plus SI effects.

### Dual effects of stretch during AP waveforms

Rhythmic cardiac AP represent even more complex voltage histories than sawtooth ramps. Lacking appropriate kinetic



FIGURE 5 Reversible, reproducible stretch changes in  $I_{mHCN2}$ . (A) Four sequential B/D/A sets.  $V_{hold} = -40$  mV, followed by the indicated steps. Gadolinium was used. The boxed trace (*bottom*) expands "run 1, D" near the X, to show unitary endogenous events. For control run, D = 0 mmHg, for runs 1–3, D = -20 mmHg. (B) Averaged traces and  $I_{diff}$  (expanded at *right*). Asterisk, nearly overlooked stretch-accelerated current decline. (C) For runs 1–3, expanded stretch (averaged D) and no-stretch (averaged (B + A)/2) segments from above show that  $I_{mHCN2inst}$  was stretch insensitive and that for ~500 ms, >1/3 of the time-dependent  $I_{mHCN2}$  was due to stretch.

data for cardiac modeling, we looked empirically at MS  $I_{mHCN2}$  during APs.  $I_{mHCN2}$  before, during, and after stretch was compared from patches clamped to cyclic AP waveforms. This was complicated by the fact that the amplitude of  $I_{mHCN2}$  changed systematically with AP frequency (it was smaller at lower frequencies) and by the fact that during low frequency runs, stretch often decreased  $I_{mHCN2}$  irreversibly. Nevertheless, when a given patch was successfully tested (i.e., when reversible stretch-induced changes were obtained) at more than one frequency, the outcome for SAN cell-like AP excursions was as is seen in Fig. 8 *A*: during a train of APs a small amount of SA occurred at high frequency (e.g., 533 bpm (beats per minute)), whereas a

larger SI occurred at low frequency (e.g., 15 bpm). Fig. 8, *B–D*, and *F* shows data for a single patch at three frequencies: 60 bpm showed SI and 240 bpm and 480 bpm showed SA. Fig. 8 *C* shows higher resolution excerpts, and Fig. 8 *D* shows their  $I_{\text{diff}}$  (overlying the voltage waveform). Fig. 8 *E* (different patch) shows three 60 bpm trials with the AP hyperpolarizing to three different extents. Reversible SI occurred in each case, with the effect accumulating over several APs then, poststretch, decaying over several APs.

The I/V relations (B/D/A; B + A averaged) for APs (e.g., Fig. 8 *F*) revealed that with the membrane potential driven by AP waveforms, mHCN2 channels furnished the membrane with a cation conductance whose degree of inward



FIGURE 6 Effect of stretch on  $I_{mHCN2}$  activation and deactivation. (A) B/D/A with early activation (scales, 4 pA, 0.1 s; *red-black bar* as in Fig. 4 A) and deactivation expanded; for ~150 ms after the hyperpolarizing step, ~1/2 the time-dependent  $I_{mHCN2}$  was SA. The -50 mm Hg  $I_{diff}$  is shown, bottom right (with its B/D/A for reference) and above, the equivalent for a smaller stretch stimulus. The prolonged (2 s) step to -40 mV showed that after a latency, stretch accelerated the current decline. The  $I_{diff}$  shapes here clarify those from truncated protocols (Figs. 3 and 5). (B and C) B/D/A traces, as explained in the text, plus an  $I_{diff}$  for which an asterisk ( $I_{cation} > 0$  pA) indicates SI  $I_{mHCN2}$ . (D) expanded  $I_{tail}$  at -40 mV for this series of B/D/A sets (labels identify the preceding hyperpolarization level). Vertical scales from top to bottom: 3 pA, 2 pA 5 pA, 5 pA; horizontal scale: 0.4 s.  $V_{hold} = -40$  mV. Lanthanum was used here and in subsequent figures.

rectification was far weaker than expected from g (V) relations implied by voltage step families or by hyperpolarizing ramp clamp (such as in Fig. 2, B and C). Wellbehaved pacemaker current is generally seen as being "on" during the diastolic potential of an AP and "off" during most of the spike (37); indeed, HCN1 behaves in this way (25). However, the outcome we noted—namely that, with or without stretch, AP waveforms keep mHCN2 channels disequilibrated at relatively high open probability throughout the AP regime—accords with recent observations of Azene et al. (25; see their Fig. 4 B) for HCN2 and HCN4.

The SAN waveforms were nearly sinusoidal, so we checked for SI/SA effects in a more asymmetrical waveform. Fig. 8 *G* shows  $I_{mHCN2}$  for a patch driven to a Purkinje fiber



FIGURE 7 Hysteresis and sawtooth ramp I<sub>mHCN2</sub> (A) Top, a hyperpolarizing then depolarizing ramp set (with a 50-ms pause at -150 mV) from +80 mV to -150 mV. A heavily filtered segment from this ramp set and from two others at higher speeds (same patch) is shown below (labels give the duration of each limb). (B) A B/D/A ramp set (D, -40 mmHg, ramps as in A,7.0 s) and its  $I_{\text{diff.}}$  (C) i. The fast-thenslow ramp protocol for the B/D/A set (D = -40 mmHg) seen in *ii* with fast (f) and slow (s) sets plotted separately alongside their respective Idiff, for which scales are 4 pA (fast) and 10 pA (slow). In iii, relative effects of stretch on rates in the two limbs are compared graphically for the slow B/D/A by normalizing (see arrows) the average no-stretch ((B + A)/2) and the stretch (D) I/V relations.

type AP (-90 to +5 mV) at 120 then at 480 bpm (*inset*). In both cases the net effect of stretch was SI behavior; at 240 bpm (not shown) stretch produced an intermediate effect in terms of the extent and rate of SI. For this waveform, reversible stretch effects were obtained for six patches tested at two or more frequencies (30–480 bpm). SI always dominated at the lower frequencies, but the frequency at which the SI/SA balance tipped had a wide range. One patch, e.g., showed SI at 60 bpm and a mixed response at 120 bpm (initial SA subsiding to SI over the nine-beat stimulus).

These findings serve as a "proof of principle" demonstration that during rhythmic activity, a given population of HCN channels can mediate both an SI and an SA cation conductance. The kinetic particulars of homotetrameric channels operating with elevated extracellular  $K^+$  at room temperature in an oocyte bilayer without auxiliary subunits are not, of course, those of native channels. For native channels in situ, therefore, the SI/SA balance might play out very differently. In diverse situations native pacemaker channels may feel bilayer stretch during voltage excursions. These include the following: the SAN during excessive filling of the right atrium, hypothalamic osmosensory neurons during lowered blood osmolarity, smooth muscle cells in distended gut, blood vessels, and bladder. What our data indicate is that under these conditions, the SI/SA balance of the pacemaker channels can be expected to vary with the waveform of the cell in question and with the duration of the stretch stimulus.



FIGURE 8 Rhythmic AP currents before during and after stretch. The data in *A*, *E*, and *G*, and *B*, *C*, *D*, and *F* are from four different patches clamped to a rhythmic SAN cell-like AP waveform (A–F) or to a Purkinje fiber-like AP waveform (*G*). All the figures except *F* (which has I/V relations for AP currents before and during stretch at the indicated beat frequencies) and *D* (in which stretch difference currents overlie the voltage waveform) are raw current traces. In *B*, the vertical arrows locate the traces shown in *C*. Vertical scales (in *B*) included for the bottom segment of each of the three current traces are 20, 10, and 10 pA for 60, 240, and 480 bpm, respectively. In *A* and *C*, before and after traces overlap. Other details are given in the text.

### DISCUSSION

### The MS responses of HCN channels

Using recombinant homotetrameric HCN channels expressed in oocytes at levels that yielded macroscopic current, we measured patch current before, during, and after membrane stretch, with pipette suction used to apply stretch. We observed no effect of stretch on the amplitude of the instantaneous component of mHCN2 current. Overall, it appeared that, as in Kv channels (15), stretch altered HCN channel gating kinetics without inducing any novel states. In Kv1 channels, the independent depolarization-driven motions of the voltage sensor accelerate with stretch (15), so for HCN channels we thought that stretch-accelerated HCN deactivation might impede hyperpolarization-induced activation of  $I_{\rm HCN}$ . But for neither of two distantly related HCN channels was that true. In both a sea urchin and a mouse isoform, stretch accelerated the activation of  $I_{\rm HCN}$ . Further character-

ization was done with the mouse isoform, mHCN2. During voltage steps, both  $I_{mHCN2}$  activation (hyperpolarization-induced) and deactivation (depolarization-induced) accelerated reversibly with stretch. Sawtooth ramp clamp (hyperpolarize-then-depolarize) revealed a voltage hysteresis that persisted with stretch.

To predict stretch responses of  $I_{mHCN2}$  during APs or to model them from our qualitative information about MS responses was not feasible. Instead, we clamped patches to rhythmic AP waveforms then applied stretch. Like others (25), we found that rhythmic AP waveforms kept HCN channels disequilibrated at a high open probability; operational I/V relations ( $I_{mHCN2}$  versus  $V_m$  during rhythmic AP clamp, e.g., +5 mV to -100 mV at 1 Hz) reflected open channel permeation at least as much as voltage-dependent gating. These I/V curves contrast starkly with those from hyperpolarizing ramp clamp (i.e., passing through 0 mV and heading to -150 mV), which elicited no current before  $\sim$ -100 mV. During APs, given patches exhibited net SI and net SA responses, depending on waveform; but overall, at lower frequencies, it was SI that dominated.

SI dominance in APs might seem surprising given the strong showing of SA effects with step and sawtooth ramp clamp. However, in those protocols, typical depolarized  $V_{\text{hold}}$  levels created a bias for the MS responses of hyperpolarization-induced transitions, which, it turned out, were SA effects. Cardiac excitability machinery (mimicked here by AP clamp) keeps V<sub>m</sub> more hyperpolarized, so our quasiphysiological waveforms had the opposite bias from that of our steps and ramps. During slow AP depolarizations (diastole), deactivation of mHCN2 channels comes increasingly into play. If, as we suspect, stretch accelerates deactivation transitions of HCN (both modes), domination by SI during slow APs seems reasonable-likewise for slow sawtooths, where SI was apparent but only after depolarization was underway. Substantial delays precede deactivation of mHCN2, perhaps explaining why SI effects accumulated over the course of several APs (and decayed in the same fashion upon stimulus release). With higher frequency APs, the interplay of dV/dt and channel kinetics evidently made mHCN2 channels hover closer to their open states. The characteristic abruptness of the net SA effects seen at high frequencies (fast onset and offset with stretch) was striking. The next section discusses which channel motions might be candidates for MS transitions.

### **Possible MS transitions**

Generically speaking, for a two-state channel, stretch-accelerated activation of current could arise from reduced stability of the closed state (accelerated opening transition) and stretchaccelerated deactivation could arise from reduced stability of an open state (accelerating closing transition). If stretch stabilized the transition state (like a catalyst, lowering the barrier between the two states), both rates would accelerate. Practically speaking, in  $I_{\rm mHCN2}$ , neither activation nor deactivation is a kinetically simple (e.g., first order) process, so we could not attribute MS responses to specific transitions. As a proxy, since HCN and Kv channels have similar voltageinduced gating motions, we consider identified MS transitions of Shaker Kv channels (Fig. 1 B). With both protein structure and gating motions being largely conserved, the forces that structure the protein-bilayer interface are likely to be broadly conserved. The unified model of Kv and HCN mode switch (22) provides an elegant framework for comparing MS responses of HCN and Kv channels.

In Fig. 1 *B*, in keeping with that framework, major HCN and Kv transitions are shown in circular four-state kinetic schemes. The two modes are labeled; mode I corresponds, in Kv, to the slow inactivated mode. The Mannikko et al. scheme (22) highlights 4 states (drawn from a 20-state model), but here, concerted transitions (two *short arrows* in mode II) are also suggested. They are depicted in mode II

Vertical arrows signify mode-switch transitions and horizontal ones (except the short open arrow), charge-displacing voltage sensor motions. Based on the left-shifted Q (V), Mannikko et al. (22) suggest HCN mode-switch motions are molecularly related to Kv slow inactivation motions. Kv activation and Kv slow inactivation accelerate to the same extent with stretch, as if the two processes have near identical S4/bilayer interactions and thus experience the same bilayer mechanical perturbation during stretch (15). Based on simultaneous gating (and/or ionic) currents and residuespecific optical measurements as well as more extensive optical scans (39-41), Ky slow inactivation (including the earliest P-inactivation step) is thought to involve lateral S4 motions, that is, a net motion in the plane of the bilayer. This makes the sensitivity of Kv slow inactivation to bilayer stretch (15) eminently reasonable. The voltage-independent stretch-sensitive motion presumably includes the initial (P type) slow-inactivation process (i.e., the mode  $II_{open} \rightarrow$ mode I<sub>inactivated</sub> switch). A question mark on the relevant mode switch transition in Fig. 1 B suggests that stretch acceleration here might explain stretch-accelerated slow inactivation in Kv channels. That picture would be even more appealing, however, if it could explain our HCN channel observations in a straightforward way, but it cannot. To explain the stretch-accelerated activation seen in HCN channels, stretch would have to accelerate opposite-going transitions, as indicated by the question mark at  $C_I \rightarrow C_{II}$  for HCN. Given the proclivity of HCN channels for "inverted gating", however, this is a possibility.

For HCN, stretch acceleration of a rate-limiting forward transition in the path modeI<sub>closed</sub> $\rightarrow$ modeII<sub>open</sub> (up arrow, Fig. 1 A) would accelerate  $I_{\rm HCN}$  activation as would stretch deceleration of a fast back-reaction. These possibilities are not mutually exclusive. As explained in the Fig. 1 B legend, asterisks indicate the class of voltage-dependent transitions identified in Kv as stretch-and-depolarization accelerated (15). HCN tail current data showed that, after a latency, stretch speeded up the depolarization-induced deactivation of  $I_{\rm mHCN2}$ . The simplest explanation for this, namely that deactivation of HCN channels (both modes) is a stretch and depolarization accelerated process, would be consistent with the Kv activation picture. The inverse asterisk in Fig. 1 B marks a voltage-dependent transition that, in Kv, decelerates with stretch (14); this last depolarization-driven motion before pore opening is a concerted step. In a Kv mutant whose concerted opening steps happen to be rate limiting (Shaker ILT), this stretch-decelerated transition yields SI activation (14), but it is not known if HCN channels open via comparable (albeit hyperpolarization-driven and thermal) concerted motions (depicted in *gray* in Fig. 1 *B*).

### Physiological and pharmacological implications of MS transitions in HCN channels

Cardiac mechanoelectric feedback phenomena are thought to require an I<sub>MScation</sub> (10,12,42). TRP-based channels are prime candidates in neurons, as they are in osmosensory neurons (43), another rhythmically active cell, but based on our observations, it will also be important to ask if the HCNbased cation channels in these cells transduce physiologically relevant force signals. During stretch and swelling, guinea pig (low intrinsic beat rate) and mouse (high intrinsic beat rate) SAN preparations make opposite responses, the former showing increased and the latter decreased rhythmicity (10). Our core finding is that HCN channels can generate both SA and SI cation current. Thus native pacemaker channels are now candidate contributors to these and similar responses. Since the SI/SA balance in our experiments produced net SI behavior at slower beat rates and net SA behavior at faster ones, it will be important to test a variety of native HCN channels in different mammalian models subjected to swelling and stretching stimuli before speculating further. The recombinant channels, of course, lacked auxiliary proteins that are likely present in situ and may tune the channels differently (44). Commenting on work by Zicha et al. (45), Giles (46) points out that during atrial flutter, the slow rate of native HCN channel deactivation manifests as what could be considered a steady-state current (e.g., as in Fig. 8 F). If, in such circumstances, the slowness of deactivation is rate limiting, stretch-accelerated deactivation might have an appreciable impact.

In addition to osmosensory neurons (which exhibit an osmosensory SI cation conductance; see references in Liu et al. (7) and Naeini et al. (43)) HCN channels are abundant in (see Table 1) baroreceptor nerve terminals, various primary afferent mechanosensory neurons, hair cells, and sperm (where, we note, spHCN expression is restricted to the highcurvature, mechanically dynamic flagellar membrane). In these locations, the mechanosensitivity of HCN channels might provide a tunable element for mechanotransduction and/or electromotility.

With two distantly related HCN channels now included among the MS voltage-gated channels, mechanosensitivity can be regarded as a family-wide trait for voltage-gated channels. The gating of all voltage-gated channels types (8,19), including HCN channels (Table 2), is susceptible to many bilayer mechanical (18) reagents. It is likely that the susceptibilities of voltage-gated channels to bilayer stretch and to such reagents are both responses to lipid stress perturbations, one being physical, the other, physicochemical. Classic models of S4-sensor motions kept the sensor sequestered in the channel protein, away from lipids, but

#### TABLE 2 Bilayer mechanical reagents and HCN channels

Propofol (GABA<sub>R</sub> antagonist)

- 1. (5  $\mu$ M) I<sub>h</sub> and I<sub>mHCN2</sub>: activation slowed, conductance decreased (58)
- 2. (5  $\mu$ M) I<sub>h</sub> and I<sub>mHCN1</sub>: activation left-shifted, conductance decreased (though not for  $I_{mHCN2}$ ) (59)
- Halothane, Isoflurane, Sevoflurane (60)Inhibition, closed state stabilization (I<sub>h</sub> and I<sub>HCN1,2</sub> activation left-shifted)

Capsazepine (61) (TRPV antagonist)

(8  $\mu$ M) Inhibits I<sub>hHCN1</sub> (activation left-shifted, slowed)

Clonidine (62) ( $\alpha$ -adrenergic antagonist)

Putative direct inhibition of Ipacemaker

Cholesterol depletion (9)

 $I_{\rm HCN4}$  activation right-shifted, deactivation slowed. Similar findings in SAN cells.

Ethanol (63)

Ih reversibly increased, activation accelerated

Sphingosine 1-phosphate (64)

 $(1 \ \mu M)$  decreased the  $\alpha$ -adrenergic-induced enhancement of I (f) Lysophosphatidylcholine (65) (a product of ischemia)

(20  $\mu$ M) I (f) amplitude decreased; activation curve left-shifted.

Most of the listed studies provide evidence that the chemicals act directly (presumably via lipid stress effects) on the HCN channels, not via other proteins such as neurotransmitter receptors.

E.g., see Bahri et al. (66) for the effects of propofol on bilayer mechanical parameters.

now, some S4 residues are known to contact bilayer lipids (17,47). The stretch and voltage dependencies of specific Kv channel transitions (14,15) constitute proof that physically induced lateral pressure profile changes alter gating rates. We suggest that the MS kinetics of mHCN2 and Kv channels follow a common pattern (Fig. 1 B) because broad conservation of structure in voltage-gated channels dictates robust conservation of bilayer mechanics. If so, modulation of voltage-gated channels by broad-spectrum lipophilic reagents may involve the same voltage sensor and gate transitions that are perturbed by membrane stretch. If this holds, much pharmacological perplexity evaporates (14,19,48). There would be no need to invoke halothane, ethanol, propofol, cholesterol, and capsazepine "binding sites" in HCN channels to explain kinetic modulation by these bilayer mechanical reagents (see Table 2) any more than we need to invoke "stretchactivation and stretch-inhibition gating motifs" in HCN channels to explain their repertoire of stretch responses.

### SUPPLEMENTARY MATERIAL

An online supplement to this article can be found by visiting BJ Online at http://www.biophysj.org.

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