Many voltage-gated potassium channels open in response to membrane depolarization and then inactivate within milliseconds. Neurons use these channels to tune their excitability. In Shaker K$^+$ channels, inactivation is caused by the cytoplasmic amino terminus, termed the inactivation gate. Despite having four such gates, inactivation is caused by the movement of a single gate into a position that occludes ion permeation. The pathway that this single inactivation gate takes into its inactivating position remains unknown. Here we show that a single gate threads through the intracellular entryway of its own subunit, but the tip of the gate has sufficient freedom to interact with all four subunits deep in the pore, and does so with equal probability. This pathway demonstrates that flexibility afforded by the inactivation peptide segment at the tip of the N-terminus is used to mediate function.
Voltage-activated potassium (K_V) channels are potassium selective integral membrane proteins formed by the assembly of four homologous subunits. In response to a membrane depolarization, K_V channels open, allowing K^+ to permeate. In many members of K_V channels, sustained depolarization leads to fast inactivation caused by an N-terminus gate. Excitable cells utilize inactivating K_V channels to shape their action potentials and adjust their firing patterns. It has been demonstrated that the inactivation gate is fully extended during inactivation such that its hydrophobic N-terminus tip interacts with residues deep in the intracellular cavity.

**Figure 1 | Effects of concatenation and I470V mutations on inactivation.**

(a) Shaker homotetramer. Cartoon represents a Shaker homotetramer channel having four inactivation gates (top) with a representative K^+ current in response to a voltage step to + 60 mV from a holding potential of − 80 mV displayed below. Red line overlaying the trace represents a fit of the inactivation process to a Markov kinetic model parameterized by the time constants k_{on} and k_{off}. Average values were: k_{on} = 430 ± 70 s⁻¹; k_{off} = 21 ± 2 s⁻¹. Horizontal bar for all panels: 10 ms; Vertical bar: 100 pA. (n = 5 patches, 43 traces). (b) Shaker concatemer having one inactivation gate. Cartoon (top) with a representative K^+ current shown below. Best fit parameter values were: k_{on} = 108 ± 21 s⁻¹ and k_{off} = 40 ± 9 s⁻¹. We noticed that k_{off} of wild-type concatemer was ∼ 2 faster than Shaker homotetramer. Vertical bar: 300 pA. (n = 7 patches, 68 traces). (c) Shaker concatemer having one inactivation gate and four subunits with I470V mutation. Cartoon (top) with a representative K^+ current shown below. Best fit parameter values were: k_{on} = 90 ± 3 s⁻¹; k_{off} = 163 ± 66 s⁻¹. Vertical bar: 100 pA. (n = 10 patches, 114 traces).

**Results**

**Construction of a Shaker K_V concatemer channel.** To study the pathway of a single inactivation gate, we constructed a Shaker concatemer channel having only one free N-terminus. Figure 1b shows a current trace from concatemer channels in response to a depolarization step. Figure 1a shows a current trace from Shaker homotetramers in response to the same depolarization step. Solid red lines overlaying the traces represent fits of a Markov kinetic model to the current traces. As expected, the extent of inactivation in the concatemers was less than that of the wild-type channels, and the relaxation slower. Quantitatively, we observed a fourfold reduction in the on rate of inactivation (k_{on}) for concatemer channels relative to Shaker homotetramers, consistent with the presence of only one inactivation gate relative to four.

**Site of inactivation gate action in the pore.** We begin by asking: does the site of action in the pore depend on the subunit to which the inactivation gate belongs? In K_V homotetramers, it has been previously established that inactivation gate's site of action is at the intracellular cavity of the channel. In particular, mutating Shaker's position 470 from isoleucine to valine (I470V) dramatically reduced the extent of N-type inactivation.

**Figure 2 | A single subunit mutated to I470V is not sufficient to reproduce the reduction in inactivation seen in the concatemer having four I470V subunits.** Cartoons of characterized mutants are shown at top, vertically aligned with their corresponding traces and K_{eq} values. Cyan colour represents a subunit mutated to I470V. Middle: representative current traces in response to a voltage step to + 60 mV from a holding potential of − 80 mV, overlaid with best-fits to a Markov kinetic model. Bottom: K_{eq} values. Horizontal bar: 10 ms; Vertical bars (from left to right): 400, 100, 500 and 100 pA. Open circles represent individual experiments and filled diamonds correspond to their averages.
Figure 3 | Multiple I470V mutations are required to modify the energetics of N-type inactivation. Top: cartoons of selected characterized mutants, where cyan colour represents a subunit mutated to I470V. Bottom: representative current traces from mutants cartooned above. All currents are in response to a voltage step to +60 mV from a holding potential of −80 mV, and overlaid with best-fits to a Markov kinetic model. Horizontal bar: 10 ms; Vertical bars (from left to right): 400, 100, 100 and 100 pA.

Figure 4 | Inactivation free energy as a function of the number of subunits mutated to I470V. Free energy was calculated from $k_{on}$ and $k_{off}$, obtained from the Markov kinetic model and plotted in RT units. The following mutants are represented in the plot: wild-type concatemer; Single I470V mutations: 1V; 2V; 3V; 4V. Two I470V mutations: 1V4V; 2V4V. All subunits with I470V mutations: 1V2V3V; 2V3V4V. The solid line represents a fit to a model in which the single inactivation gate can bind to one of four possible binding sites at the cavity. If there is a valine present, the apparent affinity of the remaining isoleucines is increased by 8% (see Methods). Data are shown in box plot format.

We reasoned that if the inactivation gate interacts with only a single and specific subunit inside the pore to produce inactivation, then a single I470V mutation at this interacting subunit would produce a reduction in inactivation comparable with that observed when all the four subunits are mutated to I470V. Current traces of Shaker concatemers with a single subunit mutated to I470V are shown in Fig. 2. No individual I470V mutation reproduced the dramatic effect observed when all the four subunits were mutated to I470V. Rather, none of the individual I470V subunit mutations produced significant reduction in the extent of inactivation as compared with the nonmutated concatemer. These results demonstrate that despite belonging to a particular subunit, a single inactivation gate does not have an exclusive site of action in the cavity. In fact, the inactivating particle samples all possible sites of action with roughly equal probability.

Given that mutating a single subunit to I470V causes no change in inactivation, we ask: what is the effect of mutating two or three subunits to I470V? Figure 3 shows representative traces from concatemer channels having one, two, three or four subunits mutated to I470V. A stepwise increase in the number of valines in the cavity produced a corresponding decrease in the extent of inactivation. If each site of action acts independently on the single inactivation gate, we would expect the free energy of inactivation (AG) to change linearly as a function of the number of subunits mutated to I470V. Figure 4 shows that this relationship is not linear; the physical principles governing this nonlinearity are unknown. The solid line represents a model in which the
Access of the inactivation gate to the pore. Having shown that a single inactivation gate has sufficient freedom to interact with all subunits deep inside of the pore, we next ask: does the gate have sufficient freedom to enter the pore through multiple intracellular entryways? It has been shown previously that the N-terminus interacts electrostatically with residues in the T1 domain, on the entryways? It has been shown previously that the N-terminus entering the intracellular cavity of the channel and the T1 domain to produce inactivation (Fig. 6b). Near the intracellular entryways, the inactivation gate may have a secondary structure with multiple polar charges causing the pathway of the gate to be spatially specific. During the ~50 ms that the inactivation gate resides in the intracellular cavity (k_off ~ 20 s⁻¹ for WT channels; Fig. 1), the gate is likely to be in an ordered state stabilized by specific hydrophobic interactions between the tip of the N-terminus and the wall of the cavity. Our data on the energetics of binding at the cavity suggest that the flexibility afforded by an unbound peptide segment at the tip of the N-terminus is used to sample all possible pathways of the gate to be spatially specific.

We conclude that the inactivation gate relies crucially on both flexibility deep in the pore and spatially specific interactions at the T1 domain to produce inactivation (Fig. 6b). Near the intracellular entryways, the inactivation gate may have a secondary structure with multiple polar charges causing the pathway of the gate to be spatially specific. During the ~50 ms that the inactivation gate resides in the intracellular cavity (k_off ~ 20 s⁻¹ for WT channels; Fig. 1), the gate is likely to be in an ordered state stabilized by specific hydrophobic interactions between the tip of the N-terminus and the wall of the cavity. Our data on the energetics of binding at the cavity suggest that the flexibility afforded by an unbound peptide segment at the tip of the N-terminus is used to sample all possible pathways of the gate to be spatially specific.

Methods

Shaker concatemer channels. All subunits were initially created in their own shuttle constructs, where unique restriction sites for concatemer construction were inserted. Subunits 1 and 2 were linked by AvrII (cyclophilin A Pro and Arg); Subunits 2 and 3 were linked by SgrA1 (cyclophilin A Pro and Arg); Subunits 3 and 4 were linked by KpnI (phosphorylating Gly and Thr). All sequences of primers used in this study are shown in Table 1. All subunits contain the following background mutations: C301S, C308S and T449V. The two cysteine mutations were originally removed to have a suitable cysteine-less channel to be used for cysteine modification experiments. These mutations have no functional consequences. T449V is a mutation that substantially reduces the extent of C-type inactivation, which greatly simplifies our analysis of N-type inactivation. Subunit 1 contains the intact wild-type N-terminus conferring the inactivation gate to the concatemer. Subunits 2, 3 and 4 are the inactivation removed versions of Shaker (ΔE–46), which completely abolish fast inactivation.

Mutagenesis. All mutations were performed using standard PCR techniques and subcloned into the GW1-CMV expression vector (Table 1). Individual subunit mutations were first subcloned into the shuttle construct, confirmed by sequencing and then inserted into the concatemer. All concatemers were subjected to a vast set of restriction digestion tests to verify that all subunits were in place and in order.
Finally, all concatemer mutants were also sequenced to verify the proper number of containing mutations.

**DNA expression.** All channel constructs’ DNA were expressed in HEK293 cells. DNA was transfected using a Nucleofector II (Amamax Biosystem) system following their recommended protocol. Experiments were performed between 1–2 days after transfection.

**Experimental solutions and electrophysiological recordings.** The intracellular solution was composed of (mM) 160 KCl, 0.5 MgCl₂, 1 EGTA and 10 HEPES (pH 7.4). The extracellular solution contained (mM) 150 NaCl, 10 KCl, 1 MgCl₂, 3 CaCl₂, 10 HEPES (pH 7.4). All chemicals were purchased from Sigma. Current recordings were obtained from inside-out excised patches, using an Axopatch 200B amplifier (Axon Instruments). Currents were sampled at 10 kHz and filtered at 2 kHz. Protocol control and sampling were done using Clampex software and a Digidata 1200 AD/DA converter (Axon Instruments). Borosilicate glass (Harvard Instrument) pipettes were pulled to about 1.5–2.5 MΩ resistance (Sutter Instrument).

**Modelling ionic currents.** Markov models were constructed with Matlab according to the following transition scheme: C₂ → C₁ → O → C₂. Simulated current was computed via \( I = \text{inP}_{\text{in}} - \text{outP}_{\text{out}} \), where \( P_{\text{in}} \) is the probability of the channel being in the open state and \( I_{\text{N}} \) is a single scale factor representing the number of channels in the patch multiplied by the unitary conductance for one channel. Traces were fit with the Levenberg–Marquardt algorithm via the Matlab program lsqnonlin. Transition rates leading to the open state were held fixed when fitting mutant with the Levenberg–Marquardt algorithm via the Matlab program lsqnonlin. Traces were fit according to the following transition scheme: C₁ → C₂ → O → C₁.

**Free energies as a function of the number of 1470V.** To model the change in free energy as a function of the number of 1470V mutations, we first determined the \( K_{\text{m}} \) associated with each subunit when all subunits were wild-type isoleucine (KeqI) and when all subunits were mutated to I470V (KeqV). We then fit our data to a model in which KeqI increases ~8% for each additional valine mutation.

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**Table 1 | Sequences of primers used in this study.**

| Primer name | Sequence |
|-------------|----------|
| PCR fwd CShWT1 (+ Avrl at C-term) | CCGTATTGCTAGCTGAGAGTACAGG |
| PCR rev CShWT1 (+ Avrl at C-term) | TATCTGTTATCTGAGAGTACAGG |
| PCR rev CShWT2 (+ Avrl at N-term) | GGTACCGCTGCTGTTGCCCTGC |
| PCR fwd CShWT2 (+ SgrAI at C-term) | CGCCGGCGAGCTGCTGTTGCCCTGC |
| PCR rev CShWT2 (+ SgrAI at C-term) | CGCCGGCGAGCTGCTGTTGCCCTGC |
| PCR rev CShWT3 (+ SgrAI at N-term) | CGCCGGCGAGCTGCTGTTGCCCTGC |
| PCR fwd CShWT3 (+ SgrAI at N-term) | CGCCGGCGAGCTGCTGTTGCCCTGC |
| PCR fwd i470V Shaker | AGCAGCTGCCGGCGCGCCGAGGTGGG |
| PCR rev i470V Shaker | GCTGACCTGCGACGTCGG |
| PCR fwd ED192,193KK | CGCCGGCGAGCTGCTGTTGCCCTGC |
| PCR rev ED192,193KK | CGCCGGCGAGCTGCTGTTGCCCTGC |

Fwd, forward; PCR, polymerase chain reaction; rev, reverse.
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Author contributions
M.H. conceived the project. G.V. and M.H. designed and performed the experiments, analysed the data and wrote the manuscript. G.V. and D.S. performed molecular biology.

Additional information
Competing financial interests: The authors declare no competing financial interest.

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