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An allosteric modulator activates BK channels by perturbing coupling between \(\text{Ca}^{2+}\) binding and pore opening

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BK type \(\text{Ca}^{2+}\)-activated \(\text{K}^+\) channels activate in response to both voltage and \(\text{Ca}^{2+}\). The membrane-spanning voltage sensor domain (VSD) activation and \(\text{Ca}^{2+}\) binding to the cytosolic tail domain (CTD) open the pore across the membrane, but the mechanisms that couple VSD activation and \(\text{Ca}^{2+}\) binding to pore opening are not clear. Here we show that a compound, BC5, identified from in silico screening, interacts with the CTD-VSD interface and specifically modulates the \(\text{Ca}^{2+}\) dependent activation mechanism. BC5 activates the channel in the absence of \(\text{Ca}^{2+}\) binding but \(\text{Ca}^{2+}\) binding inhibits BC5 effects. Thus, BC5 perturbs a pathway that couples \(\text{Ca}^{2+}\) binding to pore opening to allosterically affect both, which is further supported by atomistic simulations and mutagenesis. The results suggest that the CTD-VSD interaction makes a major contribution to the mechanism of \(\text{Ca}^{2+}\) dependent activation and is an important site for allosteric agonists to modulate BK channel activation.

BK type \(\text{Ca}^{2+}\) activated \(\text{K}^+\) channels are important in regulating neural excitability\(^1\), neurotransmitter release\(^1\), and muscle contraction\(^5,6\). Aberrant function of BK channels is associated with various human diseases such as neurological disorders\(^7\)–\(^9\), hypertension\(^10\) and impaired urinary control\(^11\). The function of BK channels relies on their activation by both elevated intracellular \(\text{Ca}^{2+}\) and depolarization of the membrane voltage. The opening of BK channels provides a negative feedback mechanism to hyperpolarize the membrane and reduce intracellular \(\text{Ca}^{2+}\) by deactivating voltage-dependent \(\text{Ca}^{2+}\) channels\(^12\). The \(\text{Ca}^{2+}\)- and voltage-dependent activation of BK channels has been an important model system for understanding allosteric mechanisms of ion channel gating\(^13\). Unlike canonical voltage gated \(\text{K}^+\) channels, such as Shaker, BK channels exhibit an intrinsic open probability \((P_0 \approx 10^{-6})\) in the absence of stimuli. \(\text{Ca}^{2+}\) binding or voltage sensor activation allosterically activates the channel by favoring the open state\(^13\). The mechanisms of voltage and \(\text{Ca}^{2+}\)-dependent BK channel activation have been described by an allosteric model\(^13\), which has successfully accounted for the functional properties of BK channel activation, pharmacology\(^13\), modulation by regulatory subunits\(^16\) and mutations\(^17,18\). The molecular bases of these allosteric activation mechanisms are not clear. Understanding these molecular mechanisms may help reveal the physiological role of BK channels, facilitate the development of therapeutic agents that modulate BK channel activation, and elucidate fundamental mechanisms of allosteric modulation.

BK channels are formed by four Slo1 subunits with structural domains that are associated with distinct functions\(^19\)–\(^21\) (Fig. 1a). In the...
membrane, a central pore is formed by the transmembrane segments S5–S6 from all four subunits. Surrounding the pore are the voltage sensor domains (VSDs) from the four subunits, comprised of the transmembrane segments S1–S4. Slo1 also contains an additional transmembrane segment S0. The cytosolic tail domain (CTD) of each Slo1 subunit harbors two Ca\(^{2+}\)-binding sites, and the four CTDs, each containing two structural motifs known as the regulator of potassium conductance (RCK), form a ring-like structure, known as the gating ring. Studies have shown that Ca\(^{2+}\) and voltage activate BK channels with distinct molecular mechanisms such that depolarization can activate the channel in the absence of Ca\(^{2+}\) binding\(^{22}\), while Ca\(^{2+}\) binding can activate the channel when the voltage sensor is at the resting state\(^{13,14,23}\). In these activation mechanisms Ca\(^{2+}\) or voltage alter the conformation of their respective sensors, and the conformational change in the VSDs is thought to be transmitted to the central pore through the transmembrane segments S5–S6.

**Fig. 1 | BC5 effects on BK channels.** a hSlo1 Channel cryo-EM structure (PDB: 6v38) docked with BC5. The membrane spanning region (S0–S6) from one subunit and the N-terminus of the cytosolic tail domain (CTD) from a neighboring subunit are amplified in the inset. The CTD and VSD are colored pink and cyan, respectively. The BC5 is colored in red and represented as molecular surface. b BC5 chemical structure. c Current traces of BK channels (control, black) and with BC5 (100 \(\mu\)M, red) at various testing voltages from −30 to 250 mV with 20 mV increment. The potential before and after testing pulses was −80 mV. Inset: tail currents after the testing pulses. d GV relationship for WT BK channels at various BC5 concentration. Solid lines are fits to the Boltzmann relation. All data were collected from independent patches, \(n = 3–12\). For detailed \(n\) for each data point, please see the “Reproducibility” section in “Methods.” e The shifts of GV relationships (red, \(V_{1/2}\): voltage where the GV relation is half maximum) and outward macroscopic current inhibition (blue, \(I_C\): control current, \(I_{BC5}\) current in BC5) depend on BC5 concentration, with \(EC_{50}\) of 2.5 ± 0.3 \(\mu\)M and \(IC_{50}\) of 1.1 ± 0.1 \(\mu\)M respectively. \(n = 3–12\), exact \(n\) for each data point is shown in the “Reproducibility” section in “Methods.” f Inhibition of the outward macroscopic current by 100 \(\mu\)M BC5 versus voltage in the presence of either 1 \(\mu\)M (filled, \(n = 3\)) or 100 \(\mu\)M (hollow, \(n = 4\)) free Ca\(^{2+}\). g Representative single-channel current recordings at 100 mV and in 100 \(\mu\)M free Ca\(^{2+}\) as control (left and black) and with 10 \(\mu\)M BC5 (right and red) from an inside out patch in symmetrical 160 mM KCl. O: open; C: closed. h Voltage dependence of single-channel current amplitudes (i-V), \(n = 3\). The data points represent the mean ± SEM for the whole paper.
change is then coupled to the opening of the activation gate via pathways of intramolecular interactions in the channel protein. Although the sensors for Ca$^{2+}$ and voltage have been identified, the allosteric coupling mechanisms between the sensors and the activation gate in BK channels are still under investigation. One of the long-standing issues is to identify the paths of intramolecular interactions within the channel protein by which cytosolic Ca$^{2+}$ binding or voltage sensor activation opens the channel pore across the membrane. Another issue is that both Ca$^{2+}$ and voltage open the same activation gate. Therefore, the respective paths for mediating voltage and Ca$^{2+}$-dependent channel opening must merge eventually. Although Ca$^{2+}$ and voltage activate BK channels via distinct mechanisms, these activation mechanisms influence each other allosterically. Where and how the allosteric paths for Ca$^{2+}$ and voltage-dependent activation merge remains a fundamental question.

In BK channels while the CTD serves as the Ca$^{2+}$ sensor, it was also shown to contribute to voltage-dependent gating, and CTD-conformation was altered by voltage-dependent activation of Ca$^{2+}$ binding at the CTD, and for the VSD to open the pore, interactions among the CTD, VSD, and the pore domains are essential. Although the structural domain interactions that are required for voltage and Ca$^{2+}$-dependent activation have been intensively studied, mechanistic roles of these domain interactions remain poorly understood. In the S10 subunit the CTD is covalently connected to the S6 by a peptide of 15 amino acids, known as the C-linker. A previous study showed that deletion or insertion of C-linker residues altered Ca$^{2+}$ and voltage-dependent activation in a manner that correlated with the C-linker length, suggesting that Ca$^{2+}$ binding to the CTD may open the channel by "tugging" the C-linker. On the other hand, the CTD also has a non-covalent interaction with the VSD of its neighboring subunit (Fig. 1a). This interaction was first revealed by the discovery that the CTD and VSD form a close interface because residues from both the CTD and VSD of the neighboring subunits form a Mg$^{2+}$-binding site, which differs from the Ca$^{2+}$ binding sites in CTD. Mg$^{2+}$ binding activates the channel by enhancing voltage-dependent activation via an electrostatic interaction with the voltage sensor. The interactions between CTD and VSD were then more clearly shown by BK channel structures. Based on the structural evidence, it was hypothesized that the CTD-VSD interaction may contribute to the coupling between Ca$^{2+}$ binding and pore opening. This notion was strongly supported by the fact that the comparison between the structure of the BK channel with Ca$^{2+}$/Mg$^{2+}$ bound and that of metal-free showed a large Ca$^{2+}$/Mg$^{2+}$-dependent change in the interface between CTD and VSD. This hypothesis is also consistent with an earlier finding that the region of the CTD located close to the VSD is important in determining the different Ca$^{2+}$ sensitivities among various BK channel homologs. Furthermore, mutations at the CTD-VSD interface were also found to alter Ca$^{2+}$ sensitivity. However, these mutations either had small effects on Ca$^{2+}$-dependent activation or, besides reducing Ca$^{2+}$-dependent activation, also altered voltage-dependent activation and resulted in an increase of the intrinsic open probability of the channel. Did these mutations alter the Ca$^{2+}$-dependent activation or did some of these mutations alter the voltage-dependent activation by affecting the voltage sensor, which then indirectly affect Ca$^{2+}$-dependent activation? Thus, whether the CTD-VSD interaction as revealed by structural data mediates coupling of Ca$^{2+}$ binding to pore opening remains unclear.

In this study we use in silico screening to identify a compound that binds to a site at the CTD–VSD interface and enhances BK channel activation. We found that the compound specifically enhances Ca$^{2+}$-dependent activation by perturbing the pathway for coupling between Ca$^{2+}$ binding and pore opening. These results support the idea that the non-covalent CTD–VSD interaction makes a major contribution to the coupling between Ca$^{2+}$ binding and pore opening that is independent of voltage-dependent activation, and that the CTD–VSD interface is an important site for allosteric agonists to enhance Ca$^{2+}$-dependent activation of BK channels.

**Results**

In order to probe if the VSD–CTD interaction of BK channels is required for Ca$^{2+}$-dependent activation, we initiated an in silico search for compounds that may interfere with BK channel Ca$^{2+}$-dependent activation. Specifically, we docked a subset of the available chemical database (Active Chemical Database (ACD, Molecular Design Ltd.) of about 4 × 10$^{6}$ compounds, in which each compound carries one or two charged groups with a net charge of 0, ±1, or ±2, to a site near the VSD–CTD interface (Fig. 1a, Supplementary Fig. 1). Patch clamp recordings of BK channels expressed in Xenopus oocytes with and without the candidate compounds from the in silico screening showed that one of the compounds, BC5 applied at the cytosolic side of the membrane patch (Fig. 1b), activated the channel as measured from the inward tail currents at a voltage (~80 mV) more negative than the equilibrium potential of the K$^{+}$ channel. Single channel recordings showed that BC5 caused brief closures during channel opening without altering the single channel conductance (Fig. 1g, h), consistent with the mechanism that BC5 acts as a pore blocker from the cytosolic side, and can be flushed out by negative voltages and inward K$^{+}$ currents, reminiscent of Ca$^{2+}$ and Mg$^{2+}$ block of the channel.

We found that the activation and inhibition of BK channels by BC5 were independent molecular processes. BC5 inhibited the channel at a lower concentration than activation, with the half-maximum inhibition concentration (IC$_{50}$) of 1.1 ± 0.1 nM (Fig. 1e, Supplementary Fig. 2b). The inhibition was dependent on voltage, with weaker inhibition at less positive potentials (Fig. 1f) and no inhibition at a negative potential (Fig. 1c). These characteristics of inhibition are consistent with the mechanism that the positively charged BC5 acts as a pore blocker from the cytosolic side, and can be flushed out by negative voltages and inward K$^{+}$ currents, reminiscent of Ca$^{2+}$ and Mg$^{2+}$ block of the channel.

Single channel recordings showed that BC5 caused brief closures during channel opening without altering the single channel conductance (Fig. 1g, h), consistent with the mechanism that BC5 acts as a discrete pore blocker with relatively fast on- and off-rates, but not fast enough to be a fast blocker that reduces single channel conductance. In addition, unlike BC5 activation, BC5 inhibition of the channel showed little dependence on Ca$^{2+}$ (Fig. 1f) or Mg$^{2+}$ (Supplementary Fig. 3a, b), and the mutations that reduced BC5 activation had little effect on BC5 inhibition (Supplementary Fig. 4a, b). These results suggest that BC5 inhibition of the channel is an off-target effect, namely, in addition to interacting with our target docking site for channel activation (Fig. 1a), BC5 may interact with the pore (Supplementary Fig. 4c) to block the outward K$^{+}$ currents.

Our target docking site of BC5 in the BK channel is located near the Mg$^{2+}$-binding site for channel activation, which is formed by residues from both the VSD and CTD. As shown in Fig. 2a, d, residue E399 is shared by both the Mg$^{2+}$- and BC5-binding sites. If BC5 interacts with the target docking site to activate the channel, we expected that an electrostatic repulsion between Mg$^{2+}$ and the charges in BC5 and a direct competition between Mg$^{2+}$ and BC5 for binding would reduce BC5 activation. Consistent with this idea, we found that BC5 at the same concentration (100 µM) induced less shift in GV relations in the presence of 10 mM Mg$^{2+}$ (Fig. 2b, c and Supplementary Fig. 3a). Likewise, a charge reversal mutation of the Mg$^{2+}$-binding residue, D99R, also reduced BC5 activation of the channel (Fig. 2c), although D99 is not part of the BC5-binding pocket, supporting that BC5 binds in the vicinity of the Mg$^{2+}$-binding site.

To further characterize BC5 interactions with the target docking site, 200 ns molecular dynamics (MD) simulations of the open and closed channel were performed to probe further the allosteric agonist BC5 binding site.
closed states of BK channel with and without docked BC5 were performed in explicit membrane and water (see "Methods") (Supplementary Table 1). The results showed multiple binding modes between BC5 and the channel that interchanged dynamically with time, and in the process BC5 interacted with different residues (Fig. 2d). As summarized in Supplementary Table 2, the hydrophobic naphthalenyl group of BC5 was buried in contact with the intracellular side of S0, S1, S3, and S4 as well as the residue F395 from αB of CTD. The charged arginine part of BC5 interacted with the negatively charged residue E399 (Fig. 2d). The binding poses of BC5 were more narrowly confined in the open conformation, suggesting that BC5 binding is stronger and may favor the open state of BK channels. We mutated the BC5 interacting residues individually or in combination to alanine or other amino acids that changed the charge of the BC5 interacting residue.

![Image of BC5 interactions with the targeted site at the CTD-VSD interface.](https://doi.org/10.1038/s41467-022-34359-6)

**Fig. 2** | BC5 interactions with the targeted site at the CTD-VSD interface. a The cryo-EM structure of hSlo1 (PDB: 6v38) with BC5 docked to its targeted site. A similar region as in Fig. 1a is shown. The VSD and CTD are colored silver and cyan, respectively. Mg²⁺-binding residues (D99, E374, E399) are represented as green sticks. The Mg²⁺ position is highlighted using the purple circle, which is hatched to indicate that Mg²⁺ and BC5 do not bind at the same time. b GV relationship of BK channels and Boltzmann fits (solid lines) in 10 mM Mg²⁺: n = 3 for control and n = 4 for 100 μM BC5. For comparison, the dashed lines represent the GV relationships of the control (black) and 100 μM BC5 (red) in the absence of Mg²⁺, taken from Fig. 1d. c GV shifts caused by 100 μM BC5. For WT the GV shifts in 0 (n = 3) and 10 mM (n = 3) Mg²⁺ are plotted, and for the D99R (n = 4) mutant, GV was measured in 0 Mg²⁺. *: p = 0.0002 and 0.0475 respectively, unpaired Student’s t-test. Data points are shown in solid circles. d Dynamic interactions of BC5 in the binding pocket of metal-free (left) and Ca²⁺-bound (right) BK channels. Snapshots of BC5 (thin sticks) were extracted from the atomistic simulation trajectories every 10 ns and superimposed onto the last frame of sim 1 and 7 (Supplementary Table 1). As a reference, representative BC5 binding poses (closed: at 180 ns from sim 7; open: at 102 ns from sim 1) are also shown in thick bonds. e GV shifts of mutant BK channels in response to BC5. n = 3–12, exact n for each data point is shown in the "Reproducibility" section in "Methods." The WT curve is taken from Fig. 1e. f Maximal G-V shift (ΔV1/2) and EC50 of BC5 for mutations and the WT. DM1: double mutation E180A/N182A; DM2: I105E/E399R. The data were obtained from fittings of the dose response in Fig. 2e using Hill equation. The standard errors of EC50 and ΔV1/2 were estimated directly from the SEMs of the input data and validated by numerical simulations.
and measured dose responses of the mutant channel activation to BC5 (Fig. 2e). These mutant channels showed increased EC_{50} (Fig. 2f), suggesting that the mutations reduced BC5 binding by interrupting the interactions between those residues and BC5. Since BC5 binds to the channel with multiple modes, a mutation to a single residue in the binding pocket may not be sufficient to abolish BC5 binding. Instead, the mutation may prevent some of the binding modes but reinforce others. This may have happened to some of the mutations, which, while enhancing EC_{50}, also increased the slope of the BC5 dose response and the maximum channel activation (Fig. 2e, f). To further disrupt BC5 binding, we made double, triple or quadruple mutations that combined these individual mutations. However, except for the two shown in Figs. 2e and 2f, the combined mutations abolished channel function. These results suggest that this pocket of residues are important for channel function. Consistently, we found that some of the individual mutations altered channel function (Supplementary Fig. 5). These results support that BC5 enhances BK channel activation by binding to the target docking site.

BK channels are activated by voltage and Ca^{2+} with distinct mechanisms. Thus, we asked whether BC5 shifted the GV relation of BK channels by modulating the voltage or Ca^{2+}-dependent activation mechanism. To address this question, we first examined whether BC5 modulates voltage activation of BK channels by measuring gating currents with and without BC5 (Fig. 3a). Unlike the GV relationship, the voltage dependence of gating charge movement (QV) was not shifted to more negative potentials (Fig. 3a). Furthermore, measurements of intrinsic openings of the channel at negative voltages (<−100 mV) when the voltage sensors were at the resting state showed that BC5 enhanced open probability (Fig. 3b). These results suggest that BC5 did not activate the channel by altering the voltage-dependent mechanism. Since the BC5- and Mg^{2+}-binding sites are located closely and share the same subunit and in a neighboring subunit. The Ca^{2+}-binding site is in the CTD, while the activation gate is located in the membrane spanning domain. In both open and closed conformations, the allostereic pathway crossed these domains via two major branches, with one branch via the non-covalent VSD-CTD interface from aA/aB in the CTD to the S4-S5 linker/S6 in the VSD, and the other branch via the covalent C-linker to S6. To complement the pathway analysis, we used the information flow between D367 (‘source’) and F315 (‘sink’), which quantifies the contributions of all the residues to dynamic coupling. The results show that the information flow through VSD-CTD interface is higher or similar to that through the C-linker/S6 connection (Supplementary Fig. 9), suggesting that this interface is critical for the coupling between Ca^{2+} binding and pore opening in addition to the C-linker/S6 pathway. Importantly, the network analysis shows that BC5 lies adjacent to the key allostereic pathways at the VSD-CTD interface, interacting with residues E395 and E399, which are part of both the BC5-binding site and the allostereic pathway in the closed structure bound with BC5 (Fig. 4a). In addition, in other closed and open structures F395 and E399 are in the vicinity of the allostereic pathway within a distance of <6 Å (Fig. 4a, b and Supplementary Table 2). These results suggest that BC5 binding likely perturbs the allostereic pathway to interact with both Ca^{2+} binding and pore opening.

We performed a mutational scanning of selected residues on or in the vicinity of the allostereic pathway and found that the mutation T245W reduced whereas E219R enhanced the sensitivity of the channel to Ca^{2+} for activation (Fig. 4c, d). T245 is part of the allostereic pathway, while E219 is in the vicinity of the allostereic pathway in the open state with or without BC5 (Fig. 4b). E219 and T245 are in S4 and S5, respectively, and are downstream from BC5 in the allostereic pathway, being closer to the activation gate (Fig. 4a, b). If BC5 perturbs the allostereic pathway to activate the channel as shown above (Fig. 4a, b; Supplementary Fig. 9) we expected that these two mutations would alter BC5 activation as well. Indeed, similar to the effects on Ca^{2+}-dependent activation, T245W reduced BC5 activation of the channel such that BC5 shifted the GV less in the mutant than in the WT channel (Fig. 4c, d), while E219R enhanced BC5 activation (Fig. 4c, d). These results support the notion that the allostereic pathway couples Ca^{2+} binding to pore opening (Fig. 4a, b) and that BC5 activates the channel.
by perturbing the pathway. Taken together, our results demonstrate the importance of the molecular interactions at the VSD-CTD interface of BK channels in coupling Ca$^{2+}$ binding to pore opening. One of the two major allosteric pathways mediating Ca$^{2+}$ binding in the CTD to open the channel gate in the membrane is via the VSD-CTD interface. It is not clear where the allosteric path eventually merges with that for voltage-dependent activation. However, since both Ca$^{2+}$ and BC5 can activate the channel independently of voltage (Fig. 3a, b) the allosteric path for Ca$^{2+}$-dependent activation starting from Ca$^{2+}$-binding sites remains distinct from voltage-dependent activation at least down to the BC5 site.

**Discussion**

We identified BC5 using in silico screening in order to modify Ca$^{2+}$-dependent activation of BK channels (Fig. 1a, b). Our results showed that BC5 activated BK channels by selectively altering the Ca$^{2+}$-dependent activation mechanism in the absence of voltage sensor activation and without indirectly modifying voltage-dependent gating.

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Fig. 3 | BC5 perturbs Ca$^{2+}$-dependent activation. a Top, gating currents in control and 100 µM BC5. Voltage pulses were from −80 to 300 mV with 20 mV increments. Bottom, Normalized gating charge-voltage (QV) relation of on-gating currents. The smooth lines are fits to the Boltzmann function, $n = 14$ for control, and $n = 7$ for BC5. b Top, current traces of a patch containing hundreds of BK channels at −140 mV in control and 100 µM BC5. Only brief unitary channel openings were seen at the negative voltage. O: open; C: closed. Bottom, Open probability (Po) at negative voltages, $n = 9$ for control, and $n = 3$ for BC5. c GV relations in various [Ca$^{2+}$]; (black hollow symbols) and in addition of 100 µM BC5 (red filled symbols). Black solid lines are Boltzmann fits without BC5 while red solid lines are Boltzmann fits with BC5. n = 3 for all conditions. d GV shifts in response to 100 µM BC5 at different [Ca$^{2+}$]. Data points are shown in solid circles, all of $n = 3$. e BC5 effects on the D367A5D5N mutant channel in 0 and 100 µM [Ca$^{2+}$]; $n = 4$ for 0 [Ca$^{2+}$] control; $n = 5$ for 0 [Ca$^{2+}$], with 100 µM BC5; $n = 4$ for 100 µM [Ca$^{2+}$] control and $n = 3$ for 100 µM [Ca$^{2+}$]; with 100 µM BC5. D367A and 5D5N (D897-901N) ablated the two Ca$^{2+}$-binding sites in each Slo1 subunit, respectively. GV relations were fit with the Boltzmann function (solid lines). f BC5 effects on the Core-MT BK channel in 0 and 100 µM [Ca$^{2+}$]; $n = 14$ for 0 [Ca$^{2+}$] control; $n = 9$ for 0 [Ca$^{2+}$] with 100 µM BC5; $n = 3$ for 100 µM [Ca$^{2+}$] control and $n = 6$ for 100 µM [Ca$^{2+}$] with 100 µM BC5. GV relations were fit with the Boltzmann function (solid lines).
Molecular docking, atomistic simulations and mutagenesis studies together suggest that BC5 binds at the CTD-VSD interface (Fig. 2), away from Ca\^{2+}-binding sites in the CTD and the pore in the membrane. Yet it not only promotes pore opening (Fig. 1d) but also interacts with Ca\^{2+} binding such that its effect on channel activation is inhibited by Ca\^{2+} binding (Fig. 3c–f). Thus, BC5 perturbs both ends of the Ca\^{2+}-dependent activation mechanism, the sensor and the pore, allosterically by interacting with the pathway that couples Ca\^{2+} binding
to pore opening. This pathway is built intrinsically in the BK channel protein, since BCS still activates the Core-MT channel (Fig. 3f), in which the entire CTD is removed and replaced with an exogenous short peptide, indicating that the pathway downstream from the BCS-binding site is preserved despite the removal of Ca²⁺ sensors.

We found that BCS activated BK channels by shifting the GV relation (Fig. 2f) and increasing open probability at negative voltages (Fig. 3d) to a similar extent as 1 mM Ca²⁺1–22. When the dependence of BK channel activation on Ca²⁺ was fit with the MWC model, the Ca²⁺-binding affinity at the open state was ~1 mM1–4. Thus, BCS activation of the channel only reached a part of the full capacity of Ca²⁺-dependent activation (Fig. 4d). This is not considered surprising because Ca²⁺ binding is coupled to pore opening via two major pathways, one is through the CTD-VSD interface, which is affected by BCS, and the other is through the C-linker19–21,27,29,30,36,38 (Fig. 4a, b). However, in our dynamic network and information flow analyses, the highest information flow residues were observed at the VSD-CTD interface (Fig. 4a, b and Supplementary Fig. 9). These results suggest that the pathway through the VSD-CTD interface may make more contributions to Ca²⁺-dependent activation than the C-linker. However, since the maximal activation by BCS was less than half of that by Ca²⁺ (Fig. 4d), BCS interaction with the pathway may not be strong enough to even achieve the full capacity of Ca²⁺-dependent activation via shifting the GV-VDSD pathway.

This idea is consistent with the result that Ca²⁺ binding inhibited BCS effects on channel activation (Fig. 3c, d), suggesting that the perturbation on the pathway by Ca²⁺ binding was stronger and overrode that by BCS.

Each Slo1 subunit of BK channels contains two Ca²⁺-binding sites in the CTD23–25,41 and we found that Ca²⁺ binding to both sites shared the same CTD-VDSD allosteric pathway for coupling to the pore. We measured Ca²⁺ effects on BCS activation of the channel with either Ca²⁺-binding site removed by mutation SDSL or D367N17. BCS activated the channel similarly to that in the WT BK in either of these mutant channels in the absence of Ca²⁺. In 100 μM Ca²⁺ BCS activation was not abolished as in the WT channel (Fig. 3c, d and Supplementary Fig. 10), indicating that the disrupted Ca²⁺ binding to either of the removed site by SDSL or D367N could no longer inhibit BCS effects via the VSD-CTD pathway. In other words, Ca²⁺ binding to both sites is required to inhibit BCS activation effects. These results indicate that both Ca²⁺ binding sites are coupled to the activation gate via the same allosteric pathway that interacts with BCS. Since BCS activated the SDSL or D367N channels to a similar extent by a similar GV shift (Supplementary Fig. 10), the two Ca²⁺-binding sites may activate the channel with a roughly equal amount via the allosteric pathway that interacts with BCS. Our dynamic pathway analysis shows that the part of the pathway downstream from the BCS site on the VSD-CTD interface to the activation gate is common to both Ca²⁺-binding sites, but the part of the pathway upstream from the BCS site to the two Ca²⁺-binding sites diverges to connect individual Ca²⁺ sites (Supplementary Fig. 11).

BCS activated BK channels as an allosteric agonist, but it also inhibited the channel as an off-target effect (Fig. 1c–e). Opposite to its activation of the channel, which was dependent on Ca²⁺ (Fig. 3c, d) but not on voltage (Fig. 3a, b), the inhibition by BCS was dependent on voltage but not Ca²⁺ (Fig. 1f). While BCS can be used as an excellent tool to study Ca²⁺-dependent activation mechanisms, it can also be used as an effective BK channel inhibitor in physiological conditions, where the membrane voltage remains more positive than the K⁺ equilibrium potential so that BK channels are always inhibited. Our results revealed the CTD-VDSD interface as an excellent site for allosteric agonists (Fig. 2) and the inner pore as a site for inhibitors (Supplementary Fig. 4) of BK channels. It should be noted that our current study suggests that BCS adopts multiple binding modes in the CTD-VDSD interface and most contacts are transient. Thus, further chemical modification of BCS may yield new compounds with more stable interactions with BK channels, which may possess stronger allosteric activation activity. These results also suggest that potentially other sites along the allosteric pathways (Fig. 4a, b) can be used for allosteric agonists or antagonists to modulate BK channel function.

These sites can be targeted for the development of therapeutic molecules to treat BK channel associated pathologies including drug therapies for BK channel associated diseases such as neurological disorders associated with either gain-of-function or loss-of-function BK mutations or impaired urinary bladder control4.5.

Methods

Molecular docking and in silico screening

A structure-based in silico screening strategy was employed in this study. This strategy was previously applied to the KCNQ1 channel, resulting in the discovery of the novel channel modulators5,6. Specifically, the in silico screening method used our molecular docking software MDock59–60 as the docking engine, and each small molecule in a chemical library was docked to a selected site on a target protein. The protein structure was treated as a rigid body during docking and other parameters in the docking program were set as default. Then, compounds in the library were ranked by their binding scores evaluated by ITScore2 (implemented in MDock)59, which is a knowledge-based atomic pairwise scoring function derived by using a statistical mechanics-based iterative method based on a training set of experimentally determined protein–ligand complex structures (a refined set of PDBbind 2012 containing 2897 protein–ligand complexes9,22).

As for the protein structure for docking, both open (Ca²⁺-bound, PDB: 6v38) and closed (Ca²⁺-free, PDB: 6v39) conformations of human BK channels were solved using Cryo-EM5. A major difference between these two conformations is the relative position between the membrane spanning region (S0-S6) and the cytosolic domain (CTD), as shown in Supplementary Fig. 1a. Because we aimed to discover chemical compounds stabilizing BK channels during their open state, the interface between the voltage sensor domain (S1-S4) and CTD was selected for the target site and the structure in the open state (PDB: 6v38) was used in the in silico screening study (see Fig. 1a and Supplementary Fig. 1b). Because the residues interacting with Mg²⁺ are located at the VSD-CTD interface and were included in our target site, the Mg²⁺ ion in the channel structure was removed before docking. The protein structure was prepared using UCSF Chimera19–21. Moreover, the experimental hit, BCS, was docked to both open and closed conformations of BK channels. The resulting binding modes were used as starting points for atomistic simulations.

Next, to prepare for the chemical library in this proof-of-concept study, considering that the VSD-CTD interface contains many charged residues (e.g., K98, D99, D173, and E219 on VSD, and R342, E374, K392, and E399 on CTD), we screened a subset of the Available Chemical Database (ACD version 1999, Molecular Design Ltd., currently is a part of BIOVIA in Dassault Systèmes, https://www.3ds.com/) in which each compound contains either one charged group (carrying a charge of +1, −1, +2 or −2) or two charged groups (with one group carrying +1 charge and one group carrying −1 charge). There were a total of ~4 × 10⁴ compounds in this chemical sub-database for screening. Because accurately predicting protein–ligand binding energies remains a challenging task for scoring functions in molecular docking programs49,50, current in silico screening methods are in pursuit of an enriched subset of potential candidates44,45. Therefore, we further visually inspected the docking modes of the top 2% compounds ranked by the scoring function, ITScore². Compounds forming strong interactions (e.g., hydrogen bonds or salt bridges) with both VSD and CTD in their docking modes were selected. One of the selected compounds, BCS, forms hydrogen bonds with N172 and D173 from VSD and S337 and E399 from CTD, respectively. At the end, 9 purchasable compounds (reported in Supplementary Table 3) were selected for experimental assays.
Mutations and expression

All mutations in this study were made by using overlap-extension polymerase chain reaction (PCR) with Pfu polymerase (Stratagene) on the template of the mbr5 splice variant of msol (Uniprot ID: Q08460)⁶. Sequencing of the PCR-amplified regions is used to confirm the mutation⁶. mRNA was synthesized in vitro from linearized cDNA using T3 polymerase kit (Ambion, Austin, TX). mRNA of mutations was injected into oocytes (stage IV-V) from female Xenopus laevis at an amount of 0.05–50 ng/oocyte. The injected oocytes were incubated at 18°C for 2–7 days before electrophysiology recordings. We have complied with all relevant ethical regulations and animal protocol is approved by IACUC at Washington University in St. Louis, USA.

Electrophysiology

All experimental data were collected from inside-out patches. The set-up included an Axopatch 200-B patch-clamp amplifier (Molecular Devices, Sunnyvale, CA), ITC-18 interface and Pulse acquisition software (HEKA Electronik, Lambrecht, Germany). Borosilicate pipettes used for inside-out patches were pulled using a Sutter P-1000 (Sutter Instrument, Novato, CA) and then wax coated and fire polished with a resistance of 0.5–1.5 MΩ.

The macroscopic currents were collected at the 50 kHz sampling rate (20 ms intervals) with low-pass-filtered at 10 kHz. A P/4 leak subtraction protocol with a holding potential of –120 mV was applied to remove capacitive transients and leak currents (except for experiments in Fig. 3b). The solutions used in recording macroscopic Ionic currents include: (1) Pipette solution (in mM): 140 potassium methanesulfonic acid, 20 HEPES, 2 KCl, 1 EGTA, and 22 mg/L nesulfonic acid, 20 HEPES, 2 KCl, 2 MgCl₂, pH 7.2. (2) The nominal 0 Ca²⁺ bath (intracellular) solution to make different [Ca²⁺]i (in mM): 140 Potassium methanesulfonate, 20 HEPES, 2 KCl, 5 EGTA, pH 7.1. (3) Basal bath (extracellular) solution to make different [Ca²⁺]i (in mM): 140 Potassium methanesulfonate, 20 HEPES, 2 KCl, 2 MgCl₂, pH 7.2.

The standard errors of EC₅₀ and Hill Coefficient were normalized to the maximum macroscopic tail current amplitudes at 80 or 120 mV. The GV relationship was determined by measuring the macroscopic current at 100 mV in the presence of 100 μM [Ca²⁺], where the single channel conductance is 273 pS and P₀ is about 1.0.

Dose response curve of G-V shifts (ΔV₅₀/Δ) or current inhibition on BCS concentration (EC₅₀) were listed below. Figure 1d, n = 12 for control, n = 3 for 0.01 and 0.03 μM BCS; n = 10 for 0.1 μM BCS; n = 4 for 0.3 and 1 μM BCS; n = 5 for 2 μM BCS; n = 6 for 5 μM BCS; n = 3 for 10 μM BCS; n = 5 for 30 and 100 μM BCS; n = 4 for 300 μM BCS.

Dose response curve of G-V shifts (ΔV₅₀/Δ) or current inhibition on BCS concentration (EC₅₀) was obtained from the fitting to the Hill equation:

\[ \theta = \frac{\Delta V_{50}}{\Delta} + \frac{1}{1 + \left( \frac{EC_{50}}{X^{n}} \right)^{1/3}} \]  

Where θ is the expected response at dosage X, \( \Delta V_{50/\Delta} \) is the maximum response for an infinite dosage, \( EC_{50} \) is the dosage at which the response is 50% of maximum, and n is the Hill Coefficient. In all of our final fittings Hill Coefficient was set to 1 since it was close to 1 during the initial fits. The curve was fitted using the “curve_fit” function Python SciPy module. The standard errors of \( EC_{50} \) and \( \Delta V_{50/\Delta} \) were estimated directly from the SEMs of the input data and validated by numerical simulations.

Reproducibility

The number of experiments (patches) n for some figure panels are listed below. Figure 1d, n = 12 for control, n = 3 for 0.01 and 0.03 μM BCS; n = 10 for 0.1 μM BCS; n = 4 for 0.3 and 1 μM BCS; n = 5 for 2 μM BCS; n = 6 for 5 μM BCS; n = 3 for 10 μM BCS; n = 5 for 30 and 100 μM BCS; n = 4 for 300 μM BCS.

Figure 1e, for GV shift, n = the same as Fig. 1d; for current inhibition by BCS, n = 12 for control, n = 4 for 0.01, 0.1 and 0.3 μM BCS; n = 5 for 1 and 2 μM BCS; n = 4 for 5 μM BCS; n = 3 for 10 μM BCS; n = 5 for 30 and 100 μM BCS; n = 4 for 300 μM BCS.

Figure 2e, for WT, n is the same as shown in Fig. 1d; for I05E, n = 4; for D173C, n = 5; for E180A, n = 6; for N182A, n = 8; for E395A, n = 5; for E399R, n = 3; for DMI (E180AN182A), n = 3; for DM2 (I05E399R), n = 3.

Atomistic simulations

The cryo-EM structures of hBK channel in the Ca²⁺-bound (PDB: 6v38) or Ca²⁺-free (PDB: 6v3g) states were used in all simulations reported in this work.⁹ Several short loops absent only in one structure were rebuilt using the other structure as template using the Swiss-PDB viewer. For example, A567-E576 and N853-E931 missing in Ca²⁺-bound structure were rebuilt using the Ca²⁺-free structure. Missing N- and C-terminal segments as well as several long loops (e.g., V53-G92, A614-V683, and D834-I870) are presumably dynamic and thus not included in the current simulations. Residues before and after the missing segments are capped with either an acetyl group (for N-terminus) or a N-methyl amide (for C-terminus). Standard protonation states under neutral pH were assigned for all titratable residues.

As summarized in Supplementary Table 1, four sets of atomistic simulations were performed: The Ca²⁺-bound state of BK channel with/without BCS (sim1-3 and 4-6, respectively) and the Ca²⁺-free state of BK channel with/without BCS (sim7-9 and 10-12, respectively). The initial binding pose of BCS was identified with docking study. In
and 10 ns for step 6) were performed, where the restrained force field and the CHARMM36m all-atom force field were used. The interaction parameters of BC5 were then assigned by identifying similar atom, bond, angle, and dihedral-angle types from similar small molecules within the CHARMM36m all-atom force field and CHARMM General Force Field (CHARMM36cgenff) (see Supplementary Fig. 12). All simulations were performed using CUDA-enabled versions of Gromacs 201874,75. The default CHARMM-GUI generated configuration was used except otherwise noted here. Electrostatic interactions were described by using the Particle Mesh Ewald (PME) algorithm76 with a cutoff of 12 Å. VdW interactions were cutoff at 12 Å with a smooth switching function starting at 10 Å. Covalent bonds to hydrogen atoms were constrained by the SHAKE algorithm77, and the MD time step was set to 2 fs. The temperature was maintained at 298 K using the Nose-Hoover thermostat78,79.

To analyze the dissociation of the receptor from the binding pocket, we performed a trajectory analysis using the Floyd-Warshall algorithm76 and the NVidia-based version of CHARMM-GUI Membrane Builder protocol71. The dissociation events were considered when the distance between the heavy atoms of the protein, BC5, and lipid was more than 10 Å. The analysis was performed using the Network View algorithm. The first 20 optimal and suboptimal paths were selected for analysis presented in this work.

**Information flow**

Information flow provides a global assessment of the contributions of all nodes to the dynamic coupling between selected “source” and “sink” nodes. This analysis complements the dynamic pathway analysis to provide additional insights on how different residues may contribute to sensor–pore coupling. For this, we first construct a network similar to the one described above. Pairwise mutual information was calculated between node i and j as follows: $I_{ij} = H_i + H_{ij} - H_j$, where $H_i$ is calculated as $\frac{1}{2} \sum_{x=1}^{N} \ln p_i(x_j)$, where $p_{ij}(x_j)$ is the fluctuation density and $x$ is the distance to the equilibrium position. Gaussian mixture model (GMM)80 is used to estimate the density. The residue network is then defined as $A_{ij} = C_{ij} M_{ij}$, in which $C_{ij}$ is the contact map. To analyze the information flow from source (S0) to sink (S1) nodes, the network Laplacian, is calculated as $L = D - A$, where $D$ is diagonal degree matrix: $D_{ij} = \sum_j A_{ij}$. The information flow through a given node (residue) is defined as $f_i = \frac{1}{2} \sum_{j} (p_i - P_i) A_{ij}$. The potentials $P$ is given by $P = \frac{L}{\lambda} b$, where $L$ is inverse reduced Laplacian and $b$ is the supply vector which corresponds to one unit of current entering at the source node that will exit at sink nodes. The magnitude of $f_i$ thus quantifies the contribution of residue $i$ to dynamic coupling between the source and sink nodes.

**Reporting summary**

Further information on research design is available in the Nature Research Reporting Summary linked to this article.

**Data availability**

The data that support this study are available from the corresponding authors upon reasonable request. Previously published accession codes used in this study: 6V38 and 6V3G. Source data underlying Figs. 1–4 and Supplementary Figs. 3–5, 9, and 10 are available as a Source data file. Source data are provided with this paper.

**Code availability**

Custom code used in this study is publicly available at GitHub (https://github.com/zhiguangjia/Analysis-code-for-BC5-Nature-communication/).

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Author contributions

Competing interests
The authors declare no competing interests.

Additional information
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