Structural basis and energy landscape for the Ca$^{2+}$-gating and calmodulation of the Kv7.2 K$^+$ channel

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The Kv7.2 (KCNQ2) channel is the principal molecular component of the slow voltage-gated, noninactivating K$^+$ M-current, a key controller of neuronal excitability. To investigate the calmodulin (CaM)-mediated Ca$^{2+}$ gating of the channel, we used NMR spectroscopy to structurally and dynamically describe the association of helices hA and hB of Kv7.2 with CaM, as a function of Ca$^{2+}$ concentration. The structures of the CaM/Kv7.2-HAB complex at two different calcification states are reported here. In the presence of a basal cytosolic Ca$^{2+}$ concentration (10–100 nM), only the N-lobe of CaM is Ca$^{2+}$-loaded and the complex (representative of the open channel) exhibits collective dynamics on the millisecond time scale toward a low-populated excited state (1.5%) that corresponds to the inactive state of the channel. In response to a chemical or electrical signal, intracellular Ca$^{2+}$ levels rise up to 1–10 pM, triggering Ca$^{2+}$ association with the C-lobe. The associated conformational rearrangement is the key biological signal that shifts populations to the closed/inactive channel. This reorientation affects the C-lobe of CaM and both helices in Kv7.2, allosterically transducing the information from the Ca$^{2+}$-binding site to the transmembrane region of the channel.

Significance

Ion channels are sophisticated proteins that exert control over a plethora of body functions. Specifically, the members of the Kv7 family are prominent components of the nervous systems, responsible for the ion fluxes that regulate the electrical signaling in neurons and cardiac myocytes. Albeit its relevance, there are still several questions, including the Ca$^{2+}$/calmodulin (CaM)-mediated gating mechanism. We found that Ca$^{2+}$ binding to CaM triggers a segmental rotation that allosterically transmits the signal from the cytosol up to the transmembrane region. NMR-derived analysis of the dynamics demonstrates that it occurs through a conformational selection mechanism. Energetically, CaM association with the channel tunes the affinities of the CaM lobes (calmodulation) so that the channel can sense the specific changes in [Ca$^{2+}$] resulting after an action potential.

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Data deposition: The NMR chemical shifts have been deposited in the BioMagResBank, www.bmrb.wisc.edu (accession nos. 34097 and 34226) and in the Protein Data Bank, www.wwpdb.org (PDB ID codes 6FEG and 6FEH).

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Results

Finding Soluble and Functional Constructs of Kv7.2-AB. The expression of the cytosolic moiety of Kv7.2 covering helices hA and hB (Fig. S1B) results in an insoluble and aggregated protein. We identified a hydrophobic region prone to be unstructured and to suffer from proteolysis and generated a construct of the Kv7.2 C terminus that encloses helices hA and hB (residues Q310–R552) with the linker deletion ΔR374–K493 (Kv7.2-hAB[Δ6L]; Fig. S1B). Bacterial coexpression of Kv7.2-hAB[Δ6L] with CaM results in a monodisperse protein complex. Experimental evidence supports the biological relevance of this construct: The interlink region between helices hA and hB is not required for channel function or assembly (19), and electrophysiological recordings of cells expressing Kv7.2-hAB[WT] and Kv7.2-hAB[Δ6L] demonstrate that the average current density and Boltzmann parameters are comparable in both channels (Fig. S1 C and D). Altogether, it can be concluded that the Kv7.2-hAB[Δ6L] construct (henceforth Kv7.2-hAB) represents a bona fide and meaningful model that retains all of the functional properties of the channel fragment.

Solution Structure of the CaM/Kv7.2-hAB Complex in the Absence of Added Ca²⁺. The solution structure of the CaM/Kv7.2-hAB complex in the absence of added Ca²⁺ was determined by NMR spectroscopy. First, chemical shift assignments were achieved using orthogonal isotopic labeling (Fig. S2) to alleviate signal overlap, and more than 95% of the ¹H, ¹³N, and ¹⁵C backbone resonances were assigned [BioMagResBank (BMRB) accession no. 34097; Fig. S2].

The Ca²⁺ association state with the CaM/Kv7.2-hAB complex was investigated using the ¹H, ¹³N chemical shift perturbations (CSPs) as reporters of the ion effect. Upon Ca²⁺ saturation, the vast majority of the CaM-perturbed residues belong to the EF hands located in the C-lobe of CaM, with no significant CSPs in the N-lobe (Fig. 1, yellow bars). Ca²⁺ depletions of the complex with EGTA produces exactly the mirror effect (Fig. 1, red bars). Thus, in the absence of added Ca²⁺, Kv7.2-hAB associates with CaM at an intermediate Ca²⁺ state in which only its N-lobe is loaded. This is consistent with the crystallographic structure of the CaM/Kv7.1-hAB complex (17), where CaM is also Ca²⁺-loaded just in the N-lobe. We will refer to this partially Ca²⁺-loaded complex as “intermediate” intCaM/Kv7.2-hAB.

The 3D structure of the intCaM/Kv7.2-hAB complex in solution [Protein Data Bank (PDB) ID code 6FEH; Fig. 2 A and B and Fig. S4] was calculated employing 5,981 NOEs, 306 dihedral angles, and 134 residual dipolar couplings (RDC) restraints (Table S1). The protein complex shows helices hA and hB wrapped by CaM (Fig. 2 A and B). A third helix composed of residues H357–R365 (hTW in Fig. 2B) is located in the flexible linker between helices hA and hB, and it does not make significant contacts with the rest of the protein complex. The pre-hA region (16, 17) is partially unstructured in intCaM/Kv7.2-hAB (Fig. 2B), but this is likely due to the channel truncation. The complex is structurally similar to the available PDB entries of CaM complexes to other Kv7 subunits (Fig. S3), showing rmds values of 1.07 Å with the CaM/Kv7.1-hAB complex (17) and 1.4 Å with the CaM/Kv7.2-hAB chimera (16), while it shows fewer similarities with the Ca²⁺-loaded CaM/Kv7.4-hAB complex (18) (rmsd = 8.1 Å, mostly due to a segmental rearrangement).

The Structural Basis of CaM/Kv7.2-hAB Ca²⁺ Gating. Ca²⁺ saturation of the CaM/Kv7.2-hAB complex results in CSPs in the Kv7.2-hAB moiety as well. To investigate the putative conformational changes as a function of Ca²⁺, we have investigated the structural properties of CaM/Kv7.2-hAB at two other concentrations of this ion. The “apo” form (devoid of Ca²⁺, apoCaM/Kv7.2-hAB) and the fully loaded Ca²⁺ complex (holoCaM/Kv7.2-hAB) were purified and found to be stable over time. The hydrodynamic radius and the stoichiometry of the complexes do not vary with [Ca²⁺], as shown by the elution profile from size exclusion chromatography (Fig. 1B), indicating that the ion-dependent structural variations are subtle, in agreement with the moderate CSPs of CaM/Kv7.2-hAB with [Ca²⁺] (Fig. S2). A structural model from apoCaM/Kv7.2-hAB was obtained based on these chemical shift changes combined with the hN-R residual dipolar couplings in an aligned medium. The resulting apoCaM/Kv7.2-hAB model (Fig. 2C) is similar to the partially Ca²⁺-loaded intCaM/Kv7.2-hAB complex, showing variations in the flexible regions (including the EF-hand loops as discussed below) and, to a lesser extent, in the hB helix interhelical orientation.

The solution structure of the holoCaM/Kv7.2-hAB complex was also determined by NMR spectroscopy (Fig. 2 D and E and Fig. S4). The assigned chemical shifts (94% of the ¹H, ¹³N, and ¹⁵C; BMRB accession no. 34226), a set of NOEs (n = 5,287), and RDCs (n = 131) were used to resolve the 3D fold of the protein complex (PDB ID code 6FEH). Even though the main structural features are maintained, the helices hA and hB have changed their relative disposition and holoCaM/Kv7.2-hAB is conformationally different from the existing structures, with rmds values of 3.3 Å, 3.7 Å, and 9.3 Å for the CaM/Kv7.1-hAB complex (17), CaM/Kv7.2-hA/Kv7.3-hB chimera (16) and CaM/Kv7.4-hAB complex (18), respectively (Fig. S3). More importantly, holoCaM/Kv7.2-hAB also significantly differs from intCaM/Kv7.2-hAB (rmsd = 3.1 Å), but the two structures can be reconciled by a segmental rearrangement (Fig. 2F): Ca²⁺ association with CaM produces a 17.9° rotation of the EF hands, which also pulls helices hA and hB of CaM and helix hA of Kv7.2. Remarkably, as a result of the segmental solid-rigid rotation, the hAB helices have changed the interhelical angle and their relative orientation with respect to CaM. Altogether, such torsion is the necessary (and sufficient) rearrangement to explain the allosteric transmission from the Ca²⁺-binding site of the CaM’s C-lobe up to the helix hA (and to the pre-hA) of Kv7.2.

All of the experiments were run at 120 mM KCl, and under these close-to-physiological conditions, the N-lobe of CaM in the complex binds Ca²⁺ with extremely high affinity. Actually, apoCaM/Kv7.2-hAB can only exist in the presence of a
large excess of a strong Ca\(^{2+}\) chelating agent (i.e., EGTA), and the Ca\(^{2+}\) affinity for the N-lobe is estimated to be subnanomolar (K\(_d\) < 10\(^{-3}\) \(\mu\)M). This is in contrast to free CaM, where the C-lobe shows the highest affinity for the ion (K\(_C\) [C-lobe] = 3.4 \(\mu\)M, K\(_C\) [N-lobe] = 14 \(\mu\)M) (20). On the other hand, the Ca\(^{2+}\) affinity for the C-lobe in \textit{intCaM/Kv7.2-hAB} has been determined by fluorescence resonance energy transfer (FRET): When two independent fluorophores are attached to Kv7.2-hAB, the above-mentioned Ca\(^{2+}\)-gated conformational rearrangement produces a change in the FRET intensity (Fig. 1C). The FRET intensity as a function of the free Ca\(^{2+}\) concentration (as determined by fura-2) (21) provides an apparent affinity constant of 0.89 \(\pm\) 0.05 \(\mu\)M, consistent with other determinations in similar contexts (22). The Hill coefficient of 1.83 suggests that the event is monitoring the saturation of CaM’s C-lobe with two ion units.

Both CaM lobes in \textit{CaM/Kv7.2-hAB} show altered affinity with respect to free CaM (calmodulation), and we focused on the conformation adopted by the EF-hand loops, since they are responsible for Ca\(^{2+}\) association. Compared with the available crystallographic (Fig. 2G) and solution (Fig. S5) structures of CaM, the N-terminal EF-hand loops of \textit{apoCaM/Kv7.2-hAB} agree better with Ca\(^{2+}\)-bound conformations. Moreover, the interhelical angle analysis (6) (Table S2) also suggests that the lobe is already in a “holo” (Ca\(^{2+}\)-bound) conformation, providing a structural explanation for the increase in affinity. However, this mechanistic explanation is constrained by the validity of the structural model of \textit{apoCaM/Kv7.2-hAB}, based on a limited number of experimental restraints (chemical shifts and RDCs). On the other hand, the EF hands of the C-lobe in \textit{intCaM/Kv7.2-hAB} have interhelical angles characteristic of a partially open conformation (Table S2), and the conformation is similar to \textit{apoCaM} structures (Fig. 2H and Fig. S5). These results are consistent with the moderate affinity increase and the micromolar K\(_d\) range observed for the C-lobe in the complex.

### The Conformational Changes Are Also Observed upon Tetramerization.

To validate whether the observed changes are also present in a tetrameric architecture, we looked for Ca\(^{2+}\)-dependent conformational changes in a channel moiety that includes the regions responsible for tetramerization (helices hC and hD of the cytosolic C-lobe of Kv7.2; Fig. S1A). The \(^{1}H\)^\(^{15}\)N–transverse relaxation optimized spectroscopy (TROSY)–heteronuclear single quantum spectroscopy (HSQC) of the CaM/Kv7.2-hABCD (Fig. S2) shows only a few peaks corresponding to residues from flexible regions, in good agreement with the spectrum of a tetrameric structure of more than 120 kDa. Instead, the methyl-TROSY spectrum (23) of the \(^{13}\)C–\(^{15}\)N methyl groups of \textit{intCaM/Kv7.2-hABCD} displays the vast majority of residues. Despite the signal overlap, the superposition of \textit{intCaM/Kv7.2-hABCD} and \textit{intCaM/Kv7.2-hABC} spectra (Fig. 3A) allows the assignment of the hAB \(\delta\)-Ile residues in \textit{intCaM/Kv7.2-hABCD}. Noticeably, the spectrum for \textit{holoCaM/Kv7.2-hABCD} displays the same Ca\(^{2+}\)-induced CSP as \textit{holoCaM/Kv7.2-hAB} (Fig. 3A), indicating that the ion-dependent conformational changes are maintained in the tetrameric complex.

FRET experiments provide further evidence that the Ca\(^{2+}\)-dependent conformational changes are also present in the tetrameric complex. Fig. 3B and C shows how FRET analysis is sensitive to the conformational change occurring in the transition between \textit{apoCaM/Kv7.2-hAB} (solid red line) and \textit{holoCaM/Kv7.2-hAB} (dashed blue line). The magnitude of such change is maintained in the \textit{intCaM/Kv7.2-hABCD} complex (Fig. 3C and green lines in Fig. 3B), while no change in the FRET signal can be detected when using a CaM mutant unable to bind Ca\(^{2+}\) (CaM[1234]; blue lines in Fig. 3B and C).

\textit{intCaM/Kv7.2-hAB} Shows Excursions to an Excited State, Compatible with \textit{holoCaM/Kv7.2-hAB}. Analysis of the conformational ensemble obtained after the experimental restraint refinement indicates

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Fig. 2. Structure of the CaM/Kv7.2-hAB complex at different calcification states. An overlay of the 10 lowest-energy conformers representing the 3D solution structure of the \textit{intCaM/Kv7.2-hAB} (A) or \textit{holoCaM/Kv7.2-hAB} (D) complex is shown. A ribbon representation of the \textit{intCaM/Kv7.2-hAB} (B) or \textit{holoCaM/Kv7.2-hAB} (E) complex shows the structural elements: CaM protein (orange); the prehelix (pre-hA), which connects segment 6 of the pore with the rest of the intracellular C terminus of the Kv7.2 channel (purple); helix hA (red); helix hTW (green); and helix hB (blue). Ca\(^{2+}\) ions are represented as spheres. The same color code is used for the ribbon representations of the \textit{apoCaM/Kv7.2-hAB} complex. (C) Ribbon representation of the structural model of the \textit{apoCaM/Kv7.2-hAB} complex. (F) Alignment of \textit{intCaM/Kv7.2-hAB} and \textit{holoCaM/Kv7.2-hAB} by the center of mass of the N-lobe of CaM displays the rotation (17.9°) of a segment of the complex upon Ca\(^{2+}\) binding to the C-lobe of CaM. The involved structural elements are h5 and hB in CaM and hA in Kv7.2-hAB. (G) Alignment of the EF-hand motifs of the N-lobe of \textit{intCaM/Kv7.2-hAB} (orange) with the equivalent motif in other reported (Ca\(^{2+}\)-loaded) CaM structures: 4RJD (purple), 4UMO (blue), 4GOW (red), and 5J03 (green), with rmsd values of 1.12 Å, 1.22 Å, 1.26 Å, and 1.39 Å, respectively. (H) Alignment of the EF-hand motifs of the C-lobe of \textit{intCaM/Kv7.2-hAB} (orange) with the equivalent motif in other reported (apo) CaM structures: 4V0C (blue), 4E50 (red), 4IQ0 (purple), and 5J03 (green), with rmsd values of 0.93 Å, 0.94 Å, 1.18 Å, and 1.25 Å, respectively.
that the intCaM/Kv7.2-hAB protein complex may undergo intermolecular flexibility. NMR is especially suitable for the characterization of protein motions at multiple time scales, and for motions in the microsecond-to-millisecond time scale, the relaxation-dispersion (RD) experiment allows full description of the exchange process, providing an estimation for the exchange rate (kex) and the population (p90) chemical shift (Δδ) of the excited state (24). Backbone amide RD experiments collected at two magnetic fields identified pervasive dynamics in CaM's C-lobe region of intCaM/Kv7.2-hAB, with measurable line broadening ranging between 1.5 and 20 Hz (Fig. 4). The individual fitting of the RD profiles shows a high degree of consistency in the obtained parameters (homogeneous kex and p90 values), so a collective fitting was attempted. Dynamic residues could be initially clustered into three different groups (groups I–III; Fig. S6) with robust statistics (Table S3). Group I entails five residues included in the construct but not corresponding to coding regions of the channel, so it is not considered further. Group II encloses 15 residues belonging to helices hA, hB, and hTW of Kv7.2. Most of the residues in group II face the C-lobe of CaM, whose dynamic residues are clustered in group III. Remarkably, group II and III residues can be interpreted using equivalent fitting parameters (Table S3), suggesting that the intCaM/Kv7.2-hAB may undergo intermolecular concerted dynamics. Groups I and III are well described by a single motional event of 1.1 ± 0.2 ms, with an excited state of 1.5 ± 0.6%. (Table S3).

The region of the complex that experiences conformational dynamics (Fig. 4) agrees well with the segmental motion upon Ca2+ association with the C-lobe of CaM/Kv7.2-hAB (Fig. 2F). Moreover, the RD-fitted Δδ values show an excellent correlation with the CSP upon Ca2+ addition to intCaM/Kv7.2-hAB (Fig. S6D). Equivalent experiments were repeated for the apo- and holoCaM/Kv7.2-hAB states. In both cases, the dynamic residues conforming the conformational exchange (dynamic residues from groups II and III) are abrogated. Thus, we conclude that intCaM/Kv7.2-hAB undergoes excursions toward an excited state compatible with holoCaM/Kv7.2-hAB. This conformational exchange is not originated from the in and out transfer of residual Ca2+ ions to the CaM-binding site since equivalent line broadening is obtained when the RD experiments are repeated in the presence of an excess of EGTA (Fig. S6G). Instead, the collective motion seems to be originated by an incomplete electrostatic cancellation from the acidic CaM and the basic Kv7.2-hAB subunits. In this hypothesis, line broadening is drastically attenuated when the ionic strength is increased (up to 500 mM KCl; Fig. S6F). This electrostatic complementarity has also been previously observed for CaM binding to its binding domain in the SK channel (25), among other systems (26).

We also investigated the dynamic properties of the δ-Ile methyl groups, which are widespread throughout the protein complex (Fig. 3A). There are three methyl groups (out of 12) showing dynamics in the intCaM/Kv7.2-hAB complex, but not in the apo- or holoCaM/Kv7.2-hAB complex (Fig. 4), indicating that side chains are equally sensing the collective dynamics of the complex. Thus, our results demonstrate that changes in Ca2+ concentration result in modulation of the dynamic behavior of the protein complex, with two resting states (apoCaM/Kv7.2-hAB and holoCaM/Kv7.2-hAB) and a highly dynamic and interconverting state (intCaM/Kv7.2-hAB).

Discussion

Ca2+ gating of K+ channels mediated by CaM is responsible for the tight control of a plethora of subcellular functions (27). However, the Ca2+ regulation mechanism for such voltage-dependent channels has remained, to the best of our knowledge, ill-defined since the first CaM-binding epitopes were identified. So far, the most accurate structural description of a Ca2+-gated regulation mechanism comes from the SK2 channels (25, 28, 29), where it has been demonstrated that Ca2+ binding to CaM produces a large conformational change that includes bridging different channel subunits. In Kv7 channels, it has been long hypothesized that Ca2+ regulation is also driven by conformational changes derived from the interaction between Ca2+ and CaM, and Minor and coworkers (18) suggested that CaM's conformational change could be the mechanical switch required to open/close the pore depending on the Ca2+ levels. In this paper, we provide extensive experimental evidence that such conformational change does indeed occur, providing an estimation of the energy landscape and a structural model for the allosteric rearrangement associated with the Ca2+ gating of the Kv7.2 channel.

Fig. 4. Millisecond dynamics of intCaM/Kv7.2-hAB populate the holoCaM/Kv7.2-hAB state to a low extent. In the structure, the spheres pinpoint the 15N-H backbone (bb) amide groups and 13C-δ-Ile methyl groups that are experiencing microsecond-to-millisecond dynamics for the three conformations under consideration. The magnitude of the exchange line broadening is color-coded as indicated. The vignettes show a representative example of the RD profiles (Val436). The solid lines correspond to the fitting to the Carver–Richards equation, simultaneously, of the two static magnetic field datasets (600 and 800 MHz).
We report the conformational instances of the CaM/Kv7.2 complex with atomic resolution at the two calcification states plus a structural model of a third state, covering all of the intracellular physiological concentrations, from 10–100 nM at the resting state up to 1–10 μM after chemical or electrical signaling: apoCaM/Kv7.2-AB (Ca$^{2+}$ unloaded), intCaM/Kv7.2-AB (loaded in the N-lobe of CaM), and holoCaM/Kv7.2-AB (full Ca$^{2+}$-loaded). In the three cases, the structures show a very intimate complex in a 1:1 stoichiometry where CaM wraps the Kv7.2 helices that are forming an antiparallel coiled-coil: Helix 6A is bound to the apo-C-lobe of CaM, whereas helix 6B is making contact with the Ca$^{2+}$-loaded N-lobe. This result is largely consistent with the crystallographic structures of the CaM/Kv7.1-hAB and CaM/Kv7.3-hA/ Kv7.2-hB complexes reported by Hirsch and coworkers (16, 17).

The reported conformational instances conceal the dynamic transitions required to define the conformational/energy landscape associated with Ca$^{2+}$ gating of the channel (energy landscape of Fig. 5 and Movie S1). The apoCaM/Kv7.2-hAB complex exists only in the presence of strong chelating agents. Moreover, apoCaM/Kv7.2-hAB results in an open complex according to in cellulo M-current recordings with a CaM variant unable to chelate the ion (CaM$^{2-33}$, Fig. 5). Since the basal Ca$^{2+}$ concentration of the neuronal cytosol is around 100 nM, we conclude that this conformation should be only theoretical and not found intracellularly. On the other hand, intCaM/Kv7.2-hAB is structurally rather similar and, energetically at least, 7 kcal·mol$^{-1}$ more stable than apoCaM/ Kv7.2-hAB (as estimated from the change in $K_d$ values), also maintaining the channel in the open state. The existence of different conformations that result in an open channel underscores that the conformational plasticity in the N-lobe may be required for putative interactions with other secondary messengers (i.e., phosphatidylinositol biphosphate) (30) to further modulate the gating of the channel.

From the functional point of view, the key step is the transition from intCaM/Kv7.2-hAB to holoCaM/Kv7.2-hAB: Ion binding to the C-lobe conversely displaces a segment of CaM (helices 6A and 6B), which is allosterically transmitted to the channel moiety by modifying the orientation of helices 6A and 6B (Fig. 2F). We hypothesize that this conformational change is mechanically transmitted to the intramembranous region of the channel through a change in orientation of the pre-hA element, gating the closing of the channel and, ultimately, resulting in M-current depletion. Such change is energetically favored at about 1 kcal·mol$^{-1}$, which ensures the reversibility of the system upon disappearance of the signaling Ca$^{2+}$ concentration.

The above-mentioned states are not isolated conformations only connected by the (increasing) Ca$^{2+}$ concentration; rather, they are also intrinsically coupled through thermal motions. The RD experiments show pervasive dynamics in CaM's C-lobe of intCaM/ Kv7.2-hAB, which can be adequately described as a collective motion of the complex. In the most simple and widespread interpretation of the RD experiment, the two-state model chemical shift analysis demonstrates that intCaM/Kv7.2-hAB is sensing a low-populated state (1.5% at 303 K), which corresponds to the holoCaM/Kv7.2- hAB state. Such dynamics are likely originated by an insufficient structural complementarity between the (acidic) CaM and the (basic) Kv7.2-hAB in intCaM/Kv7.2-hAB, which becomes fully satisfied (and the millisecond dynamics largely quenched) when an excess of Ca$^{2+}$ locks the complex in the holoCaM/Kv7.2-hAB state. From the functional point of view, the preexisting equilibrium of populations is a consequence of the (low) energy barrier between the states, and Ca$^{2+}$ association with intCaM/Kv7.2-hAB implies a conformational selection mechanism instead of a more canonically induced fit. Conformational selection is widely used by nature, and has been reported in many enzymes (31) and binding proteins (32), including CaM (33), and it is most efficient to produce an allosteric response after a stimulus (34).

The results presented here are also able to explain the role of CaM and the calmodulation mechanism associated with the Ca$^{2+}$ gating of the Kv7.2 channel (Fig. 5). At any ion concentration, the Kv7.2-hAB complex subunit cannot be purified in the absence of CaM. No aggregation is observed in the absence of CaM (Fig. 5A) and the affinity for the channel is so high that the complex cannot be dissociated by any of the in vitro-test conditions. Likely, this is also the case in vivo, due to the high intracellular concentration of CaM (10$^{-5}$–10$^{-5}$ M) (5). Together, these results suggest that CaM association with Kv7.2 may become consubstantial to the channel's integrity once both molecules interact. This situation is similar to the homologous channel Kv7.1, where a constitutive binding role was hypothesized (17), and to other proteins like phosphorylate kinase (5). In any case, CaM association with Kv7.2 results in a significant alteration of the Ca$^{2+}$ affinities of the two lobes, sensitizing the CaM-effector complex to the specific [Ca$^{2+}$] in the different neuronal states. At the structural level, such changes in affinity are consistent with the conformational changes observed in the EF hands of CaM/Kv7.2-hAB, compared with free CaM (Fig. 2F and G and Table S2). The N-lobe of free CaM has a low affinity for calcium (20), which is drastically increased upon association with Kv7.2 ($K_d < 10^{-3}$ μM in apoCaM/Kv7.2-hAB, Fig. 5). Considering that the basal Ca$^{2+}$ concentration in the neuron is around 100 nM, the N-lobe should always be Ca$^{2+}$-loaded. On the other hand, the C-lobe of CaM also alters its affinity for Ca$^{2+}$ upon binding to Kv7.2 (Fig. 5), lowering its apparent $K_d$ from 3.4 μM (for free CaM and at this ionic strength) (20) down to 0.9 μM in the intCaM/Kv7.2-hAB complex. This affinity tuning is required for the complex to rest inactive until an agonist-induced increase in the intracellular Ca$^{2+}$ concentration occurs (up to 1–10 μM; Fig. 5). The higher [Ca$^{2+}$] shifts the equilibrium toward the holoCaM/Kv7.2-hAB conformation, with the subsequent
conformational change that is allosterically transmitted up to the pre-helix element and likely to the transmembrane region, ultimately resulting in the closing of the channel.

Finally, it is worth considering the implications that channel tetramerization may have on the mechanisms introduced here. The methyl-TROSY spectra of the CaM/Kv7.2-hABCD complex demonstrate that the same conformational states are maintained after channel tetramerization of the C terminus (Fig. 3A), while FRET experiments prove that Ca\(^{2+}\) triggers a conformational change in the tetrameric protein. According to our model, four CaM units will bind the cytosolic moiety of a full channel, generated as previously described (20).

Methods

Expression and Purification of the CaM/Kv7.2-hAB Complex. The Kv7.2-hAB segment (residues 316–532) with the deletion ΔK374-K493 (Δk) was purified as previously described (20).

NMR Spectroscopy and Structure Calculation. NMR data were collected at 303 K on an 800-MHz Bruker Avance III spectrometer equipped with a cryoprobe and a 120-MHz Bruker Avance III US2 spectrometer. Resonance assignments were obtained by the combined use of \(^{1}H\)-HSQC, \(^{1}H\)-COSY, HNCA, HNCOa, HNCOb, HNCOc, HNCCoA, HNCaH, HNHaA, HnCh, HnCaH, HnCCoN, NOE connectivities were obtained using \(^{1}H\)-NOESY- HSQC and \(^{1}H\)-NOESY-HSQC (120-ms mixing time, 16-rs rotation correlation time). Residual dipolar coupling was extracted by using TROSY \(^{1}H\)-HSQC and semi-TROSY \(^{1}H\)-HSQC experiments in isotropic and anisotropic (C125F1-hexanol) conditions. Protein complex structure was determined using Ari3a (33). Transversal RD experiments were always acquired at two fields (600 MHz and 800 MHz) using a relaxation compensated-pull Carr-Purcell-Meiboom-Gill sequence (CPMG) sequence (80 μs of total CPMG time) and variable effective fields: 25, 50(±2), 75, 100, 125, 150, 200, 250, 325, 400(±2), 500, 600, 700, 800, 900, and 1,000 Hz.

Electrophysiological Recordings. For the WT constructs, all experiments were carried out using HEK293T cells (HEK 293T/17, CRL-11268; American Type Culture Collection). The data were acquired at a sampling rate of 1 kHz and filtered at 100 Hz, and were analyzed using pCLAMP software (version 8.2; Axon Instruments) and plotted using SigmaPlot.

FRET Experiments. All FRET experiments were carried out using an AMINCON Bowman series 2 luminence fluorimeter. The fluorescence emission spectra of the proteins collected at 500 nm were collected over 470–570 nm after exciting at 458 nm (4-nm bandwidth). The FRET index was calculated as the ratio between the emission at 485–490 nm for the blue donor and the emission at 520–525 nm for the yellow acceptor. The total protein in each condition was assessed by direct excitation of the yellow protein at 520 nm, collecting the emission at 520–570 nm.

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