Calmodulin and Ca\(^{2+}\) control of voltage gated Na\(^{+}\) channels

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Abbreviations: Nav, Voltage activated sodium channels; CIP, channel interacting proteins; CaM, calmodulin; Cav, voltage activated calcium channels; CTNav, C-terminal Nav; CDI, calcium dependant inactivation; CaM C-lobe, C-terminal lobe of calmodulin; CaM N-lobe, N-terminal lobe of calmodulin; EFL, EF hand like domain; FHF, fibroblast growth factor homologous factor; rmsd, root mean square deviation.

The structures of the cytosolic portion of voltage activated sodium channels (CTNav) in complexes with calmodulin and other effectors in the presence and the absence of calcium provide information about the mechanisms by which these effectors regulate channel activity. The most studied of these complexes, those of Nav1.2 and Nav1.5, show details of the conformations and the specific contacts that are involved in channel regulation. Another voltage activated sodium channel, Nav1.4, shows significant calcium dependent inactivation, while its homolog Nav1.5 does not. The available structures shed light on the possible localization of the elements responsible for this effect. Mutations in the genes of these 3 Nav channels are associated with several disease conditions: Nav1.2, neurological conditions; Nav1.4, syndromes involving skeletal muscle; and Nav1.5, cardiac arrhythmias. Many of these disease-specific mutations are located at the interfaces involving CTNav and its effectors.

Voltage activated sodium channels (Nav) are the major participants in the depolarization and electrical conduction of neurons, muscle cells and heart cells. They consist of a major subunit (\(\alpha\)) whose activity is modulated by several channel interacting proteins (CIP). The \(\alpha\) subunit contains 4 pseudo-repeats, each containing 6 transmembrane helices which form the Na\(^{+}\) conducting pore and the voltage sensor, plus a C-terminal region that includes an EF-hand-like motif (EFL) and a long helix (helix \(\alpha VI\)) followed by a region of variable length depending on the isoform. The different isoforms (Nav1.1 to Nav1.9 and Na\(_{\alpha}\)) also differ in their tissue distribution, cellular localization and the specific effectors that control their function. Recent publications have provided structural insight into the regulation of several sodium channels (Nav1.2, Nav1.4, and Nav1.5) by 2 of the most important effectors: calmodulin—a CIP—and Ca\(^{2+}\). Another CIP, the fibroblast growth factor homologous factor (FHF, such as FGF11, FGF12, FGF13 or FGF14), which has an inhibitory effect on channel activity, is present in several relevant structures. The cytoplasmic C-terminal is the region of the channels that interacts with these effectors. This region is also the locus of many disease causing mutations.

Mutations in the genes coding for these 3 Nav channels have been associated with neurological conditions (Nav1.2; autism\(^5\), infantile spasms and bitemporal glucose hypermetabolism\(^6\), febrile seizures\(^7,8\)), with skeletal muscle disorders (Nav1.4; myotonia\(^9-11\) and paralysis\(^12,13\)), and cardiac arrhythmias (Nav1.5; Long QT syndromes\(^14\), Brugada syndrome\(^15,16\), cardiac conduction defect, atrial fibrillation) and dilated cardiomyopathy\(^14\).

Here we review and discuss the regulation of these channels by calmodulin (CaM) and Ca\(^{2+}\) with emphasis on the specific structural characteristics of the C-terminal region that may be responsible for their differences in functional behavior of the isoforms.

Comparison of the Nav1.5 and Nav1.2 structures in their CaM complexes

The structure of the Nav1.5 C-terminal fragment in complex with CaM (CTNav1.5-CaM) was determined in the presence of Mg\(^{2+}\) (PDB ID 4OVN) and CTNav1.2 was crystallized in complex with CaM and FGF-13 in the presence of Ca\(^{2+}\) (PDB ID 4JPZ). These 2 structures show a high conservation of secondary structure elements, as expected from their high sequence similarity (91% sequence similarity, 80% identity) (Fig. 1A). However, despite the high sequence homology, there are differences between the 2 structures as shown by the main chain trace of Nav1.5 (PDB ID 4OVN) with the tube radius representing the mean r.m.s. deviation of the \(\alpha\)-carbons of the equivalent residues of the 2 chains\(^3,4\) (Fig. 1B). Most of the differences in Ca\(_{\alpha}\) positions result from a change in the orientation of helix \(\alpha VI\) with respect to the EFL between the Nav1.5 and the Nav1.2 structures (distance between the ends of the helices of ~14 Å; see Figure 2). A similar distance is observed when the CTNav in the
The structure of the CTNav1.5-CaM complex is aligned with that in the Nav1.5-CaM-FHF (Fig. 2).

Residues involved in the interface EFL-FHF, in particular the loops between αII and αIII, αIII and β2, and loop L4 and strand β4, are conserved between Nav1.5 and Nav1.2 suggesting that FHF may play a similar role in the regulation of the long-term inactivation of both isoforms by interacting with the EFL of CTNav (Fig. 1C, D). 2,3

Interaction of Nav with calmodulin

In complexes involving Nav, CaM lobes have been observed in all 3 conformations described previously: open, semi-open, and closed. 17 CaM has 2 lobes (N- and C-lobe) each containing 2 EF-hands both of which can bind Ca2+. Each lobe can, in principle, adopt any of the 3 conformations. The open conformation is observed when Ca2+ is bound to the EF-hands, while the other 2 are observed in the structurally equivalent apoCaM and Mg2+ bound CaM (Table 1).

The semi-open conformation of the CaM C-lobe in the CTNav1.5-CaM structure (PDB ID 4OVN) is similar to that of apo CaM bound to Nav peptides such as the apo CaM C-lobe complex with the Nav1.2 IQ motif peptide, (PDB ID 2KXW; rmsd 1.0 Å; Fig. 3). 18 Although the CTNav1.5-CaM-Ca-FHF and the CTNav1.5-CaM-Ca-FHF are reported as Ca2+ bound, the arrangement of the CaM helices in the C-lobe does not resemble the open conformation typically described in bound CaM-Ca2+ structures such as those observed in Cav-CaM-Ca structures. 19,20 It is noteworthy that the authors indicate that the

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**Figure 1.** Sequence and structural similarity of Nav1.5 and Nav1.2 in their cytoplasmic C-terminal region. (A) Red background displays sequence identity, yellow background sequence conservation of charge and white background, non-homologous residues. The relative accessibility (acc), calculated by Endscript/DSSP, is shown in white for buried residues and in blue for accessible residues. 19 Residues involved in the intermolecular contact between 2 Nav1.5 molecules are labeled “H” (contacts with distances less 3.2 Å, black H; those with distances between 3.2 and 5.0 Å, red H). Helix αII interacts with the CaM N-lobe and helix αVI with the CaM C-lobe (D; colored as for the “H”). Residues at the end of helix αVI involved in intermolecular contact between two Nav1.5 molecules in CTNav1.5-CaM (or with CaM N-lobe in CTNav1.5-CaM-Ca-FHF) are labeled “G.” The end of helix αVI interacts with another molecule of Nav1.5. The light gray boxes show the residues of Nav1.2 that interact with FHF in the ternary complex (PDB ID 4JPZ). (B) Co trace of CTNav1.5, in which the thickness of the tube is proportional to the rmsd of the Ca residues in the structural overlap of CTNav1.5 (PDB ID 4OVN) with the CTNav1.2 (PDB ID 4JPZ); (non-conserved residues in gray). (C) As (B) with CaM bound as in 4OVN. (D) Surface representation of the Nav1.5 showing complete sequence conservation at the surface that binds FHF (green); Nav1.5 is 180 degrees from the orientation in (B and C).
Figure 2. Snapshots of Nav1.5 structures aligned by their EFL domain. (A) Heterodimer of the structure of Nav1.5-CaM. (B) Nav1.5 bound to FHF and with CaM in extended conformation. (C) Nav1.5 bound to FHF with CaCaM as in 4JQ0. In (A–C), the structures are aligned by the EFL domain. (D) Close-up of helix αVI displaying the position of K1922, R1910 and K1899. (E) Similar to panel (B) but showing the same side chains as (D) to display the rotation of 90° around the helix αVI axis relative to the position of the other Nav1.5 molecule (light gray). (F) Displacement of helix αVI of the CTNav1.5-CaM-FGF (yellow) with respect to the CTNav1.5-CaM-FHF (green).

Table 1. Available CTNav structures. The table includes the residues of CTNav used in the study, the residues and lobe of CaM, whether or not Ca²⁺ was present in the experimental conditions, the orientation of CaM with respect to CTNav and the PDB ID of the structure. All the structures listes have the CaM C-lobe bound to the Nav IQ motif in a semi-open conformation.

| Identifier       | Amino Acids | FGF   | CaM  | Ca²⁺  | N-lobe Config. | N-lobe Target | CaM Orient. | Method   | PDB ID |
|------------------|-------------|-------|------|-------|----------------|---------------|-------------|----------|--------|
| IQNav1.5-CaM     | 1901–1927   | no    | 1–148| apo   | closed         | none          | NA          | NMR      | 2L53¹⁷   |
| IQNav1.2-CaM     | 1901–1927   | no    | 76–148⁸ | apo  | NA           | NA            | NA          | NMR      | 2XXW¹⁸   |
| CTNav1.5-CaM-FGF | 1773–1940  | yes   | 1–148| apo   | closed         | none          | NA          | X-ray    | 4DCK²    |
| CTNav1.5-CaM-Ca-FGF | 1773–1940 | yes   | 1–148| Ca²⁺  | open          | post-IQ       | anti-parallel | X-ray    | 4JQ0³    |
| CTNav1.2-CaM-Ca-FGF | 1777–1937 | yes   | 1–148| Ca²⁺  | open          | post-IQ       | anti-parallel | X-ray    | 4JPZ³    |
| CTNav1.5-CaM     | 1773–1929   | no    | 1–148| apo   | closed         | EFL           | parallel     | X-ray    | 4OVN⁴    |

*Hs: Homo sapiens, unless indicated.
*Pt: Paramecium tetraurelia

CaM N-lobe is not bound, resulting in no determination of CaM lobes' orientation relative to Nav.

¹Only the CaM C-lobe was used, resulting in no determination of CaM lobes' orientation relative to Nav.
²The authors called the N-lobe in 4DCK semi-open, and closed in 4JPZ.³ See text and Fig 4 for discussion.
anomalous signal of Ca\(^{2+}\) in the C-lobe was weak. Given the weakness of the anomalous signal and the fact that the data extend only to 3.8 Å, it is possible that a majority of the CaM C-lobe molecules do not contain Ca\(^{2+}\). Alternatively, the strong interaction of the C-lobe with helix \(\alpha\)VI may prevent the lobe from adopting the open conformation even in the presence of Ca\(^{2+}\).

The N-terminal lobe of CaM (CaM N-lobe) is observed in different conformations in the complexes with CTNav: closed, and open (Fig. 4).\(^{17}\) The structural similarity of the CaM N-lobe in the structure of the CTNav1.5-CaM-FHF (PDB ID 4DCK) and the structure of the isolated CaM N-lobe without Ca\(^{2+}\) (rmsd 1.0 Å with PDB ID 3UCW; Fig. 4A, B) is in agreement with, that as reported, this structure does not contain Ca\(^{2+}\) (rmsd ~2.7 Å with 1CDM, a complex with Ca\(^{2+}\); Fig. 4C, D). Comparison of the alignment of CTNav1.5-CaM N-lobe with CaM N-lobe-Mg (rmsd 0.4 Å, Fig. 4E, K) vs with CaM-Ca (rmsd 4.1 Å, Fig. 4H, N) emphasizes the lack of Ca\(^{2+}\), and the presence of Mg\(^{2+}\) in the CTNav1.5-CaM structure. On the other hand, alignment of the CaM N-lobe with Ca\(^{2+}\) to either CTNav1.5-CaM-FHF (rmsd 1.4 Å, Fig. 4I, O) or CTNav1.2-Ca-CaM-FHF (rmsd 0.8 Å, Fig. 4J, P) compared to the alignment of the same 2 structures with the CaM N-lobe-Mg (rmsd 3.4 Å Fig. 4F, I; and rmsd 3.2 Å Fig. 4G, M) agrees with the presence of Ca\(^{2+}\) in the N-lobe of both structures (Fig. 4F–P).

The C-terminal lobe of apoCaM (CaM C-lobe) interacts with the Ile-Gln of the IQ motif of helix \(\alpha\)VI of the C-terminus of all Nav channels. This interaction seems to serve as an anchor for the control of activation of the channels by CaM (Fig. 3). The apoCaM C-lobe recognition of the CTNav helix \(\alpha\)VI shows structural characteristics typical of the complexes of CaM with the helices of the large number of other CaM regulated proteins (Fig. 5).\(^{18,20-25}\) The residues that participate in this interaction in the case of Nav have been identified in several structures.\(^{2,4,18,21,26}\) These residues are mostly conserved in Nav1.2, Nav1.4, and Nav1.5 (Fig. 1A and Fig. 6). In addition to the isoleucine and the glutamine of the IQ motif, several residues in the middle of helix \(\alpha\)VI are also conserved (Fig. 1 and Fig. 6). Of the 14 residues of helix \(\alpha\)VI in contact with the CaM C-lobe, 10 are conserved and 4 are similar (Fig. 1A and Fig. 6), underscoring the equivalence of the interactions between the helix \(\alpha\)VI of the 3 Nav channels and the CaM C-lobe.

In the structure of the CTNav1.5-CaM complex, the N-terminal lobe of CaM (CaM N-lobe) interacts with the CTNav1.5 EFL (Fig. 1C, 2A).\(^{4}\) This interaction involves contacts of helices \(\alpha\)1 and \(\alpha\)2 of the CaM N-lobe with helix \(\alpha\)III of the EFL (residues 1832–1838) and the beginning of helix \(\alpha\)VI (residues 1895–1899) of the CTNav1.5 (Fig. 1A, 2A). Binding of FHF is not compatible with this interaction (see below). The sequences of both Nav1.5 regions involved in contacting the CaM N-lobe are conserved in the 3 Nav channels considered in this review. The helix \(\alpha\)III residues that contact the CaM N-lobe in the Nav1.5-CaM complex (QISLI) have sequences in Nav1.2 (KVQLI) and Nav1.4 (KIKLI) that appear to be compatible with the same EFL/CaM N-lobe interaction. The sequences of

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**Figure 3.** CTNav1.2 and CTNav1.5 aligned by their CaM C-lobe. This alignment displays the different contacts of the CaM N-lobe: parallel in CTNav1.5-CaM (olive) and antiparallel in CTNav1.2-CaM-Ca-FHF (marine). The alignment shows that the EFL domains do not align due to the 90° degree rotation of helix \(\alpha\)VI. The FHF domains have been omitted for clarity. (A) Alignment of CTNav1.5-CaM with CTNav1.5-CaM-Ca-FHF. (B) CTNav1.5-CaM (CTNav1.5, purple; CaM, marine). (C) CTNav1.5-CaM-Ca-FHF (CTNav1.5, yellow; CaM, olive). (D) CTNav1.2-CaM-Ca-FHF (CTNav1.2, orange; CaM, teal).
the other contact region involving the N-terminal of helix αVI (TL(R/K)RK) is highly conserved among the 3 proteins. These observations suggest that the interaction of the CaM N-lobe with the CTNav is a common feature of the 3 Nav channels (Fig. 6).

The structure of the CTNav1.5-CaM complex can be described as a "parallel" arrangement to reflect the fact that the N-lobe of CaM is located toward the N-terminal EFL domain of CTNav1.5 (Fig. 2A, B and 3A, B). On the other hand the CTNav1.5-CaM-Ca-FHF is an "antiparallel" arrangement with the N-lobe of CaM located toward the C-terminus of CTNav1.5 (Fig. 3C, D). This change in orientation of the CaM N lobe has been explained by its binding of Ca2+ that appears to favor post IQ binding.2 In the structure of CTNav1.5-CaM-FHF, determined in the absence of Ca2+ by the same group, the N-lobe of CaM does not contact Nav1.5 (Fig. 2B).

In summary, residues observed interacting with CaM in the structure of the CTNav1.5-CaM are conserved in the 3 Nav channels with a small number of exceptions, some participating in contacts of the parallel arrangement with the CaM N-lobe [1832-1834, 1837, 1900 (Fig. 3 and Fig. 6)], and some of the antiparallel arrangement [1915, 1918–1920 (Fig. 1, Fig. 3C and D)].

**Interaction of Nav with another molecule of Nav: Nav-Nav dimerization**

An intermolecular Nav1.5-Nav1.5 interaction was observed in the structure of the CTNav1.5-CaM complex between the EFL domain of one Nav1.5 molecule (helices αI and αV) and the end of helix αVI of another CTNav1.5. Residues involved in this interaction are conserved between Nav1.5 and Nav1.2 (marked with H in Fig. 1A) suggesting that this CTNav-CTNav interaction may be have a similar functional role in both channels. Noteworthy is the fact that the only EFL residues that differ between Nav1.5 and Nav1.2 at this CTNav-CTNav interface are in contact with each other in the CTNav1.5-CaM crystal (S1920 and A1882 in Nav1.5, are K1147 and N1109 in Nav1.2; Fig. 2).

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**Figure 4.** CaM N-lobe conformations in Nav-CaM complexes. The structures of the CaM N-lobe in the different complexes are compared with the isolated CaM N-lobe with Mg2+ (green, 3UCW40,41) or CaM (magenta, 1CDM40). Helix αVI is shown in the structures in which there is an interaction between the CaM N-lobe and the helix. In the figure, CTNav1.5-CaM-FHF (navy blue, PDB ID 4DCK), CTNav1.5-CaM (marine, PDB ID 40VN), CTNav1.5-CaM-Ca-FHF (olive, PDB ID 4JQ0), CTNav1.2-CaM-Ca-FHF (teal, PDB ID 4JPZ), and CaM-Ca (magenta, PDB ID 1CDM) are shown. (A) Overlap of the CaM N-lobe as seen in CaM N-lobe with Mg2+ (PDB ID 3UCW) with CTNav1.5-CaM-FHF (navy). (B) 90° rotation from A. (C) Overlap of the CaM N-lobe as seen in CaM-Ca (PDB ID 1CDM) with CTNav1.5-CaM-FHF (navy). (D) 90° rotation from C. (E, K) Alignment of the CTNav1.5-CaM N-lobe with CaM N-lobe-Mg (rmsd 0.4 Å). (H, N) Alignment of the CTNav1.5-CaM-Ca-FHF N-lobe with CaM N-lobe-Mg (rmsd 3.4 Å). (I, O) Same as (F, L) with CaM-Ca (rmsd 1.4 Å). (G, M) Alignment of the CTNav1.2-CaM-Ca-FHF N-lobe with CaM N-lobe-Mg (rmsd 3.2 Å). (J, P) Same as (G, M) with CaM-Ca (rmsd 1.45 Å).
end of helix αVI, displays less sequence identity between the 2 channels, the conservation of the EFL residues suggests that the intermolecular CTNav-CTNav interaction may be functional in both Nav channels, even though this interaction has not been observed in the published Nav1.2 structures.3,18

Conformations of CTNav in the structures of CTNav with CaM and FHF

The binary complex of CTNav1.5 with CaM (CTNav1.5-CaM) and the ternary with FHF (CTNav1.5-CaM-FHF) provide snapshots of the states of the cytoplasmic C-terminal domain during the cycle of opening and closing the channel. Structural alignment of the EFL domain of the 3 known structures shows that helix αVI is not in the same orientation with respect to the CTNav EFL in all the structures: in the structures containing FHF helix αVI is rotated 90° about its long axis (Fig. 2). A change in the conformation of residues 1892–1895 (Loop L4), which link the EFL to helix αVI allows the 90° rotation of the helix. As a result, the positively charged residues of helix αVI that interact with the EFL (helix αV) of the other Nav1.5 molecule (Arg1910, Lys1899 and Arg1922) are rotated, most likely preventing or disrupting this interaction (Fig. 2D–F). The change of conformation in loop L4 which results in the 90° rotation of helix VI has been proposed to reflect a change from the inactivated state to the state that is poised for activation.4

In the structure of the CTNav1.5-CaM-Ca-FHF complex, helix αVI forms a different angle with the CTNav EFL than in the ternary CTNav1.5-CaM-FHF complex (Fig. 2F). This difference has not been suggested to be associated with a unique physiological state of the channel.

Residues involved in the EFL-FHF interface observed in the CTNav1.5-CaM-Ca-FHF structures, in particular strand β4 and the preceding loop L4, are conserved between Nav1.5 and Nav1.2 suggesting a similar role for FHF in the regulation of both channels (Fig. 1C).2,3

Figure 5. CaM C-lobe conformations bound to Nav C-termini. The structures display a semi-open, not the open conformation typical of Ca-CaM. In the figure the colors are CaM C-lobe (yellow, PDB ID 2KXW), CTNav1.5-CaM (marine, PDB ID 40VN), CTNav1.5-CaM-FHF (navy blue, PDB ID 4DCK), CTNav1.2-CaM-Ca-FHF (teal, PDB ID 4JPZ), IQNav1.2-apoCaM (yellow, PDB ID 2KXX), and CaM-Ca (Magenta, PDB ID 1CDM). (A) Overlap of the semi-open CaM C-lobe (PDB ID 2KXW) as seen IQNav1.2-CaM with CTNav1.5-CaM-FHF (PDB ID 4DCK). (B) 90° rotation from A. (C) Overlap of CaM C-lobe as seen in CaM-Ca (PDB ID 1CDM) with CTNav1.5-CaM-FHF (PDB ID 4DCK). (D) 90° rotation from C. (G-K) and (M-P) Alignment of the CTNav1.5-CaM-Ca-FHF C-lobe with CaM-Ca C-lobe (rmsd for all pairwise alignments is 1.0 Å). (E, H) Alignment of the CTNav1.5-CaM-Ca-FHF C-lobe with CaM-Ca-Ca-FHF C-lobe ( rmsd 1.3 Å). (F, I) Alignment of the CTNav1.5-CaM-Ca-FHF C-lobe with CaM-Ca-Ca-FHF C-lobe (rmsd 1.3 Å). (G, J) Alignment of the CTNav1.5-CaM-Ca-FHF C-lobe with CaM-Ca-Ca-FHF C-lobe (rmsd 1.3 Å). (K, L) Alignment of the CTNav1.5-CaM-Ca-FHF C-lobe with CaM-Ca-Ca-FHF C-lobe (rmsd 1.3 Å). (M, N) Alignment of the CTNav1.5-CaM-Ca-FHF C-lobe with CaM-Ca-Ca-FHF C-lobe (rmsd 1.3 Å).
Ca\(^{2+}\) dependent inactivation

Calcium dependent inactivation (CDI) in the presence of CaM is an essential property of voltage activated Ca\(^{2+}\) channels (Cav). Recently, it was shown that the same phenomenon is exhibited by Nav1.4.\(^{2,7}\) Surprisingly, Nav1.5 does not show this inactivation. Figure 6 highlights the location of the residues that are different between the Nav1.4 and Nav1.5 channels. In the portion of the Nav cytoplasmic region that is present in the CTNav1.5-CaM structure there are a small number of sequence differences between Nav1.4 and Nav1.5: the major differences are beyond the end of helix aVI. If the structural elements that contribute to CDI are located in the C-terminal of the channels, they must include residues that differ between Nav1.5 and Nav1.4 to account for their differences in CDI. Since CDI is likely to involve CaM, at least part of the effect must reside in the residues that differ between the 2 channels in the portion of the channels that interact with CaM—i.e., the residues showed in Figure 6.
Nav mutations and disease

Mutations in SCN5A, the gene that encodes the Nav1.5 channel, have been found in patients with LQT3 or Brugada Syndrome.14,15,28,29 The mutations in the CT include changes at positions 1774, 1779, 1784, 1787, 1795, 1803, 1825, 1839, 1895, 1901, 1903, 1909, 1911, 1913, 1919, 1924 (LQT3; Fig. 7) and positions 1774, 1795, 1812, 1861, 1872, 1901, 1904, 1919, 1924 (Brugada Syndrome).30-37 Some of these mutations occur at the CTNav1.5-CaM-C-lobe interface (1901, 1909), the CTNav1.5-FHF (1839 LQT3), or close to this interface (1839, LQT3; 1872, Brugada Syndrome). Other mutations affect the interaction of one CTNav1.5 with another CTNav1.5 (1790, 1795 and 1913 in LQT3; 1787, 1919 and 1924 in Brugada Syndrome).

Mutations in the gene coding for Nav1.4 (SCN4A) result in myotonias and other syndromes involving skeletal muscles. Residue Gln1633 of Nav1.4 is equivalent to residue 1807 of Nav1.5, also a Gln (Fig. 6). Assuming similar roles for this residue in both proteins, the mutation Gln1633 of Nav1.4 would introduce a charge at the interface between 2 Nav channels that may be incompatible with maintaining this contact. The same interface is affected by the F1705I mutation (F1879 of Nav1.5; Figures 1, 6). Similarly, if FHF does interact with Nav1.4, the E1702K mutation of Nav1.4 (Glu1876 in Nav1.5, Fig. 6) would interfere with this interaction. Interestingly, these mutations occur in residues that are conserved among the 3 isoforms.

Mutations in the SCN2A gene, which encodes for the neuronal channel Nav1.2, underlie seizure and autism spectrum disorders.6,7,38 Mutation D1793N, observed in epilepsy patients, is located in helix α1 of the EFL domain that interacts with the CaM N-lobe (Fig. 8).8 Arg1918, which interacts with the linker of the N-lobe the C-lobe when Ca$^{2+}$ is bound to CaM, was observed mutated to His in a patient with febrile seizures.7 In the CTNav1.2-CaM-Ca-FHF

![Figure 7](https://example.com/figure7.png)

**Figure 7.** Location of mutations of Nav1.5 observed in Brugada Syndrome and LQT3. (A) Surface representation of one CTNav1.5 monomer (purple) interacting with another CTNav1.5 (green). CaM is indicated only in the foreground CTNav1.5. This state was postulated to be poised for activation. (B) Brugada Syndrome mutations are at the CTNav1.5-CaM-C-lobe interface (colored as is A). CTNav1.5 mutations present in LQT3 (1774, 1779, 1784, 1787, 1825, 1839, 1895, 1901, 1903, 1924) are shown in gray and Brugada Syndrome mutations (1774, 1812, 1826, 1837, 1861, 1795, 1919, 1924) in navy blue. (C) Inset showing the FHF to display the mutations that are at the interface of the EFL domain of CTNav1.5 and the FHF (1839 LQT3) or close to it (1840, LQT3; 1872, Brugada Syndrome). (D) Close-up of the interaction of CTNav1.5-CTNav1.5 (1795 and 1913, gray, LQT3; 1919 and 1924, blue, Brugada Syndrome).

![Figure 8](https://example.com/figure8.png)

**Figure 8.** Location of mutations of Nav1.2 observed in autism and febrile seizures. (A) Surface representation of CTNav1.2 (top view, orange) interacting with CaM (teal) and FHF (lime green). Mutation of residue 1793 of the EFL domain is shown in gray. (B) CTNav1.2 mutations in gray (1902, 1918). (C) Close-up of residue R1902 close to the interface of CTNav1.2 with FHF. (D) Close-up of the position of residues Asp79 and Asp81 of CaM in the CTNav1.2-CaM-Ca-FHF (antiparallel; teal) vs CTNav1.5-CaM (parallel; marine).
complex, Arg1918 interacts with CaM residues Asp79 and Asp81. This interaction may act as a sensor of the presence of Ca$^{2+}$ in CaM: compared with the structure of CTNav1.5-CaM in the absence of Ca$^{2+}$ this portion of the linker is the pivot for positioning the CaM N-lobe. In particular, CaM Asp79 is in the same position and CaM Asp81 is 7 A away (Fig. 8D). The mutation R1902C seems to impair Ca$^{2+}$ binding 38 to the CaM in the complex, but its position in the Ca$^{2+}$ bound structure does not support this proposal (Fig. 8).

Concluding Remarks

Atomic resolution structural information is accumulating on the conformations of the cytoplasmic C-terminal fragments of Nav channels and CaM lobes in Nav-CaM and Nav-CaM-FHF complexes in the presence and absence of Ca$^{2+}$. Some of these structures have been proposed to be associated with specific functional states of the channels. The structures contain direct information about the specific interactions of the C-terminal domain of the channels that govern the control of channel activity by C2Ps and by Ca$^{2+}$. As a consequence they are providing insights about the mechanisms of altered channel function by mutations associated with the phenotypically variant sodium channelopathies. Further experimentation, including electrophysiological measurements, binding experiments, and determination of additional structures will be essential for the identification of all the states of the cytoplasmic components of the Nav channels needed to provide a full understanding of their regulation of channel activity.

Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

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