Single-molecule localization of Na_v1.5 reveals different modes of reorganization at cardiomyocyte membrane domains

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ABSTRACT

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Mutations in the gene encoding the sodium channel Na_v1.5 cause various cardiac arrhythmias. This variety may arise from different determinants of Na_v1.5 expression between cardiomyocyte domains. At the lateral membrane and T-tubules, Na_v1.5 localization and function remain insufficiently characterized. We used novel singlemolecule localization microscopy (SMLM) and modeling to define nanoscale features of Na_v1.5 localization and distribution at the lateral membrane, groove, and T-tubules in wild-type, dystrophin-deficient (mdx) mice, and mice expressing C-terminally truncated Na_v1.5 (Δ SIV). We show that Na_v1.5 organizes as distinct clusters in the groove and T-tubules which density and distribution partially depend on SIV and dystrophin. We found that overall reduction in Na_v1.5 expression in mdx and ΔSIV cells results in a non-uniform redistribution with Na_v1.5 being specifically reduced at the groove of Δ SIV and increased in T-tubules of *mdx* cardiomyocytes. Na_v1.5 mutations may therefore site-specifically affect Na_v1.5 localization and

distribution depending on site-specific interacting proteins.

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INTRODUCTION

Proper function of the heart heavily relies on the cardiac sodium channel Na_v1.5, Na_v1.5 generates the rapid upstroke of the cardiac action potential; thus, it is pivotal for cardiac excitability, conduction, and muscle contraction^{1,2}. Mutations in its gene SCN5A have been associated with many cardiac arrhythmias, such as long-QT syndrome type 3, Brugada syndrome, and sick-sinus syndrome³. Mutations can cause loss or gain of function of Na_v1.5, but the wide diversity in cardiac arrhythmias remains unexplained. The key may lie in the distribution of Na_v1.5 over specific membrane domains – "pools" – of the cardiomyocyte. Na_v1.5 pools are regulated by different proteins, so a given Na_v1.5 mutation may affect a specific pool if it disturbs the interaction with a specific protein. Na_v1.5 expression at the intercalated disc⁴ and at the lateral membrane⁵ is well established (**Figure 1**) while a third pool at the T-tubules is still disputed⁶. At the intercalated disc, where two cardiomyocytes are tightly coupled electrically and mechanically, Na_v1.5 interacts with ankyrin-G, connexin-43, and N-cadherin, among others⁴. At the lateral membrane, where costameres connect the cytoskeleton to the extracellular matrix (Figure 1a,b), Na_v1.5 interacts with the syntrophin/dystrophin complex, which is important for Na_v1.5 stability, and CASK, among others⁷⁻⁹. The lateral membrane is shaped distinctly with valley-like grooves and hilly crests¹⁰, and Na_v1.5 is expressed at both locations in murine cardiomyocytes (**Figure 1a**)^{11,12}. T-tubules are deep invaginations of the lateral membrane that originate in the groove. They mainly provide a large surface for calcium handling, and are crucial for the uniform contraction of ventricular cardiomyocytes¹³. Consequently, the majority of the voltage-gated calcium channels Ca_v1.2 is expressed at the T-tubules^{14,15}. The claim of a T-tubular Na_v1.5 pool has so far only been supported by inconclusive results. By comparing whole-cell I_{Na} densities in detubulated and control cardiomyocytes, T-tubular sodium channels were determined to conduct 30% of total sodium current¹⁶. This is probably a considerable overestimation as this method relies on the membrane capacitance as a measure for cell membrane area, but underestimates the surface area of cardiomyocytes by about 50% ¹⁷. Therefore, the presence of a T-tubular Na_v1.5 pool remains undetermined.

Our current state of knowledge regarding the localization and distribution of $Na_v1.5$ at the lateral membrane and T-tubules is based on standard microscopy methods that are limited in resolution and sensitivity. At the T-tubules, previously published immunofluorescence data on $Na_v1.5$ show an intracellular striated pattern for $Na_v1.5$, but without a T-tubular marker, this pattern cannot be attributed to the T-tubules^{18,19}. The respective confocal microscopy techniques moreover lacked resolution to resolve the narrow T-tubular structure^{18,19}. At the lateral membrane, cell-attached and scanning ion conductance microscopy (SICM) recordings of the sodium current have provided information on the number of functional channels in a given area at the lateral membrane or specifically at the crest and groove^{11,12,20,21}, but they did not provide the molecular features needed to characterize $Na_v1.5$ organization at these domains. Moreover, previous studies did not correlate the signal of $Na_v1.5$ with other structural markers such as α -actinin for the groove and Bin1 for the T-tubules^{18,19,22}. As such, current models of the roles of

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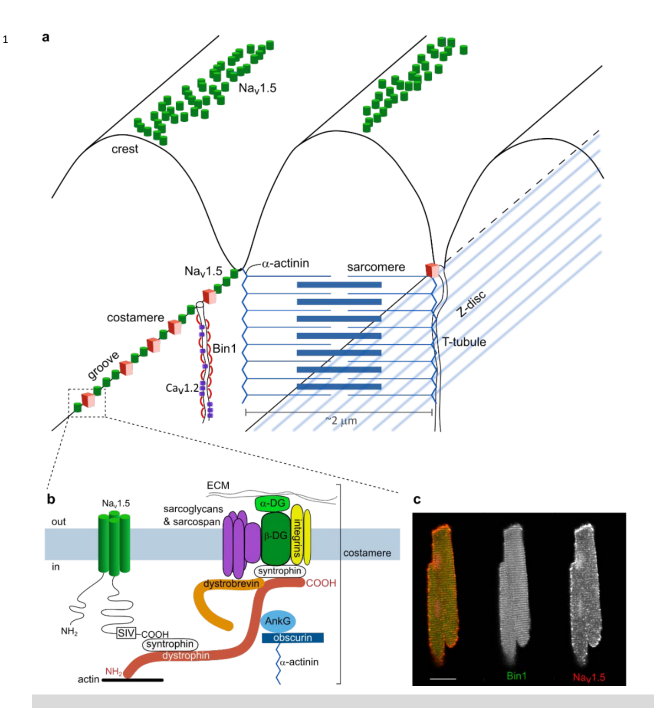


Figure 1. Pools of Na_v1.5 in cardiomyocytes. a, The lateral membrane has a characteristic profile: at the groove, the membrane is anchored to the sarcomeric Z-line (α -actinin; blue) through costameres (red boxes). T-tubules follow Z-lines and open to the groove. Their membrane is shaped by Bin1 (red curves) and contain, among many other proteins, voltage-gated calcium channels (Ca_v1.2; purple). The crest has a dome shape. Na_v1.5 is expressed at the crest and the groove. The amount of Na_v1.5 in this image serves illustrative purposes only and does not reflect experimental values. b, At the groove, Na_v1.5 (green; left) is associated with the costamere (right). The C-terminal SIV motif of Na_v1.5 binds syntrophin (white), which binds dystrophin (red). Dystrophin also binds syntrophin associated with the costamere, which contains multiple transmembrane proteins, including integrins (yellow), sarcoglycans (purple), and β-dystroglycan (green). β-dystroglycan connects the sarcolemma with the extracellular matrix (ECM) *via* α-dystroglycan. Dystrophin also links the costamere to the cytoskeleton through its association with actin at its N-terminus and with α-actinin via ankyrin-G and obscurin. c, Confocal image of murine ventricular cardiomyocyte stained for Na_v1.5 (red) and T-tubular marker Bin1 (green), showing Na_v1.5 expression at the intercalated disc, lateral membrane, and an intracellular punctate pattern coinciding with Bin1 signal. Scale bar 20 μm.

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Na_v1.5 at the lateral membrane and T-tubules do not represent the true molecular nature of Nav1.5 expression and organization at these domains. To address these issues, we utilize multi-color single-molecule localization microscopy (SMLM)²³, a super-resolution imaging method, providing the features of Na_v1.5 organization and related markers with nanoscale resolution. Using this approach, we addressed Na_v1.5 expression and cluster organization at the T-tubules using the T-tubular marker Bin1²⁴, and Na_v1.5 affinity for the groove using the groove marker α -actinin in isolated cardiomyocytes. We also assessed T-tubular I_{Na} by comparing whole-cell $I_{\rm Na.}$, rather than cluster densities, from control and detubulated cardiomyocytes. We then addressed the consequences of dystrophin deficiency (mdx) and Na_v1.5 C-terminal truncation (\Delta SIV) on Na_v1.5 expression and organization at the lateral membrane overall, at the groove specifically, and at the T-tubules. Δ SIV resembles the p.V2016M mutation found in a Brugada syndrome patient, replacing the C-terminal valine with a methionine, which results in an apparent reduced expression of Na_v1.5⁵. ΔSIV knock-in mice display a loss of sodium current and Na_v1.5 expression specifically at the lateral membrane⁵, and interaction of Na_v1.5- Δ SIV with the syntrophin/dystrophin complex is impaired⁵. In accordance, dystrophin-deficient mdx mice display a strong reduction of sodium current and Na_v1.5 expression, both overall and specifically at the lateral membrane^{7,8}. By characterizing Na_v1.5 organization and expression at different pools at a nanoscale and addressing their pool-

specific determinants, we come closer to understanding the pathophysiological variety in patients carrying SCN5A mutations, as mutations may affect Na_v1.5 pools and shape the pathophysiological landscape differently. The two genetic mouse models investigated in this study give insight into pool-specific effects of a Na_v1.5 mutation (Δ SIV) and of disturbing Na_v1.5 regulation (mdx). It appears that (1) Na_v1.5 is expressed in T-tubules; (2) dystrophin deficiency increases T-tubular Na_v1.5 expression; and (3) Na_v1.5 expression is reduced at the lateral membrane of mdx and Δ SIV mice overall whereas groove expression is specifically reduced in Δ SIV cells; (4) Na_v1.5 cluster organization changes at the lateral membrane of mdx cells and in the T-tubules of mdx and Δ SIV cells compared to wild type.

25 RESULTS

NA_V1.5 OCCURS AT DISTINCT POOLS IN MURINE VENTRICULAR CARDIOMYOCYTES

The main focus of this work is the characterization of $Na_v1.5$ localization and cluster organization in different cardiomyocyte domains with novel quantitative single-molecule localization microscopy (SMLM) and modeling. Technical validation of SMLM is shown in **Supplementary Figure 1**. Firstly, however, we gained a general overview over the $Na_v1.5$ pools with confocal microscopy. To this end, we immunostained isolated murine ventricular cardiomyocytes for $Na_v1.5$ and the T-tubular marker Bin1. $Na_v1.5$ is clearly expressed at the lateral membrane and intercalated disc (**Figure 1c**). Some $Na_v1.5$ seems to co-localize with Bin1, the latter showing a striated pattern that indicates the T-tubular membrane. The limited resolution and sensitivity of the

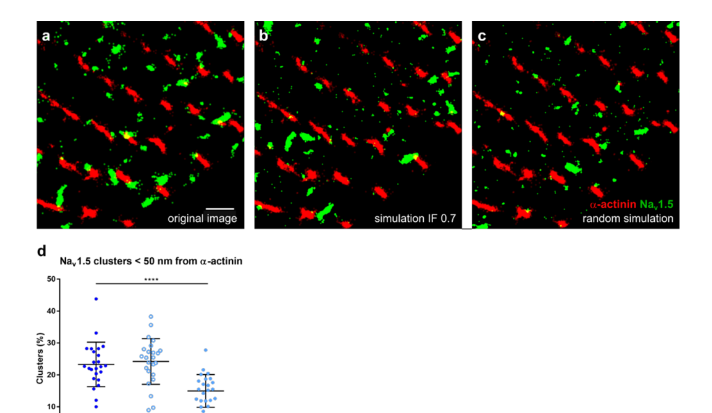


Figure 2. Na_v1.5 has affinity for the groove at the lateral membrane of a cardiomyocyte. a, Super-resolution image of the lateral membrane of murine ventricular myocyte stained for Na_v1.5 (green) and α-actinin (red). Na_v1.5 occurs close to α-actinin, indicating the groove; and in between α-actinin lines, indicating the crest. Scale bar 1 μm. b and c display simulations of the original SMLM image (a), in which Na_v1.5 clusters were randomly distributed (b) or had a high affinity for α-actinin (interaction factor [IF] 0.7 [c]). Note that differences in affinity are hard to spot by eye, especially in a small area of a cell. d, In original SMLM images of wild-type cells (N = 3, N = 24), the proportion of Na_v1.5 within 50 nm of α-actinin is ~30% higher than in simulations in which Na_v1.5 clusters of each image are distributed randomly, and similar to images in which a high affinity of Na_v1.5 for α-actinin was simulated (IF 0.7). *****, p < 0.0001, Mann-Whitney test.

- ${\small 1} \qquad confocal \ microscope \ however \ prevents \ us \ from \ establishing \ a \ detailed \ quantitative \ analysis \ of \ Na_v1.5 \ in \ the \ T-top \$
 - tubules (**Figure 1c**).

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Simulations

To determine the molecular characteristics of Na_v1.5 expression at the lateral membrane by SMLM, we imaged the lateral membrane of wild-type cardiomyocytes stained for Na_v1.5 and α -actinin, which coincides with the groove (N = 3, n = 24; **Figure 2a**). Indeed, Na_v1.5 occurred both in close proximity to α -actinin and in between α -actinin clusters, indicating localization at the groove and the crest, respectively (**Figure 1a**, **Figure 2a**). To quantify whether the Na_v1.5 clusters have an affinity for the groove, we used a novel analysis approach²⁵ based on Monte-Carlo simulations for each SMLM image (N = 3, n = 24) in which Na_v1.5 clusters were redistributed over the image either randomly or with a high affinity for α -actinin (**Figure 2b,c**). The random simulations were performed by selecting the Na_v1.5 clusters and stochastically redistributing them within the boundaries of the

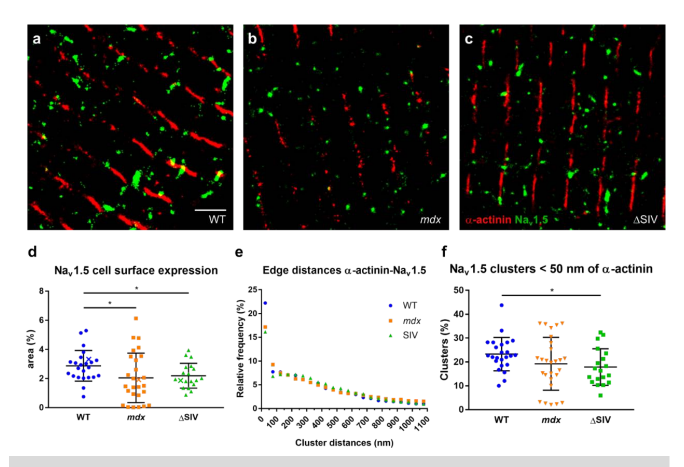


Figure 3. ΔSIV and mdx cardiomyocytes show loss of Na_v1.5 expression at the lateral membrane. a-c, Detail of SMLM images of the lateral membrane from wild-type (WT; a), mdx (b), and Δ SIV (c) cells. Scale bar 1.5 μm. d, Total Na_v1.5 expression at the lateral membrane is reduced by ~30% in mdx and Δ SIV compared to wild type. Data points represented by a cross (X) correspond to examples from panels a-c. e, Frequency distribution plot of edge distances from any α -actinin cluster to the closest Na_v1.5 cluster, bin size 50 nm. In mdx and Δ SIV, a smaller proportion of Na_v1.5 resides within 50 nm to α -actinin compared to wild type. Frequency distributions for cluster distances larger than 50 nm are similar between the genotypes. f, Groove expression of Na_v1.5, defined as Na_v1.5 clusters within 50 nm of Bin1, is reduced by ~30% in Δ SIV cardiomyocytes compared to wild type. Note that mdx cells display high variability in Na_v1.5 cluster expression at the lateral membrane (d) and at the groove (f). Wild type N = 3, n = 24; mdx N = 3, n = 27; Δ SIV N = 3, n = 19. *, p = 0.024 (wt vs. mdx [d]); p = 0.016 (wt vs. Δ SIV [d]); p = 0.019 (wt vs. Δ SIV [f]), Mann-Whitney test.

user-defined region of interest (ROI). For high-affinity simulations, an interaction factor of 0.7 was assumed, given that an interaction factor of 0.0 represents random overlap between the two colors, an interaction factor of 1.0 perfect overlap, and an interaction factor equal to or more than 0.5 indicates that one color has significant affinity for the other color²⁵. To generate high-affinity simulations with an interaction factor of 0.7, each $Na_v1.5$ cluster was first placed randomly within the region of interest, then the program determined whether one or more pixels of the cluster overlapped with an α -actinin cluster. If this overlap was detected, the cluster was kept in place; if not, the program generated a random number between 0 and 1. If that number was greater than 0.7, the cluster was kept in its position; if not, this process was repeated. Overall, these high-affinity simulations generated an image in which $Na_v1.5$ clusters overlapped with α -actinin clusters to more than a random degree.

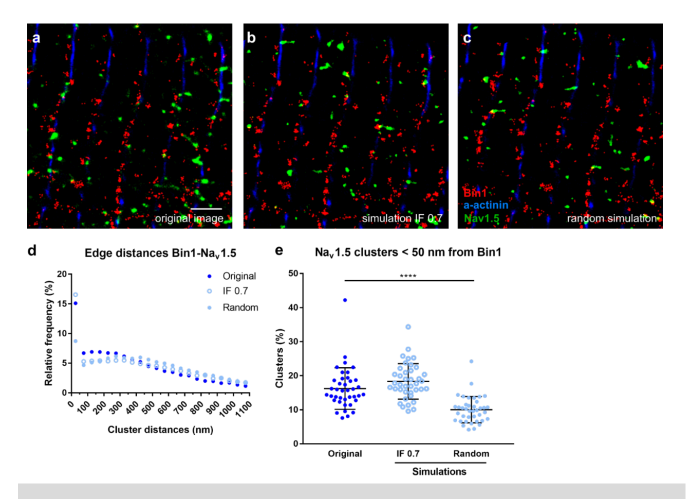


Figure 4. Na_v1.5 is expressed in the T-tubules. a, Detail of an intracellular recording from a wild-type cardiomyocyte stained for α -actinin (blue), Bin1 (red), and Na_v1.5 (green). All Na_v1.5 clusters are redistributed over the image either randomly (b) or with a high affinity for Bin1 (interaction factor [IF] 0.7 [c]). Scale bar 1.5 µm. d, For each Bin1 cluster, the closest Na_v1.5 cluster is within 50 nm in ~15% of cases, which is significantly more than the ~8% from random simulations and close to ~17% of the high-affinity simulations (IF 0.7) (N = 3, n = 39). The frequency distributions of Na_v1.5 for distances to α -actinin of more than 50 nm does not show any apparent difference between the genotypes. e, In experimental SMLM images, a higher proportion (~40%) of Na_v1.5 are within 50 nm from the nearest Bin1 cluster than in random simulations, but not as much as in simulations with a high-affinity simulations (IF 0.7). Values for each individual cell/simulation are plotted. *****, p < 0.0001, unpaired t-test.

- Then, for the original and simulated images, the distance from the edge of each α -actinin cluster to the edge of the closest Na_v1.5 cluster was determined. Na_v1.5 was considered to be expressed in the groove if it was located within 50 nm from an α -actinin cluster. We found that 25% of Na_v1.5 clusters were in the groove in experimental images. This value was close to that of the simulations with high affinity for α -actinin (**Figure 2d**), whereas in random simulations, only 15% of Na_v1.5 clusters were within 50 nm from α -actinin. These results strongly suggest that Na_v1.5 has an affinity for the groove.
- NAv1.5 EXPRESSION AT THE LATERAL MEMBRANE AND THE GROOVE DEPENDS ON DYSTROPHIN AND THE
- 8 ΔSIV MOTIF, WHEREAS CLUSTER ORGANIZATION DEPENDS ON DYSTROPHIN ONLY

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- 1 Our SMLM-derived metrics indicate that Na_v1.5 cluster density at the lateral membrane is reduced by about
- 2 30% in mdx and Δ SIV cardiomyocytes compared to wild-type cells (**Figure 3a-d**; wild type N = 3, n = 24; mdx
- N = 3, N = 27; $\Delta SIV N = 3$, N = 19). Cell size and average $Na_v 1.5$ cluster size did not change between the
- genotypes, although mdx cells showed a high variability in Na_v1.5 cluster size (**Supplementary Figure 2a**).
- Interestingly, the solidity and circularity of $Na_v 1.5$ clusters was markedly increased in mdx cells compared to
- 6 wild type (Supplementary Figure 2a). Given that solidity is defined as the ratio between the particle area and
- the convex hull area of the particle, and circularity as the roundness of a particle (1.0 indicating a perfect circle),
- 8 the increased solidity and circularity together indicate that Na_v1.5 clusters are of geometrically less complex
- shapes. The overall reduction of Na_v1.5 expression at the lateral membrane in \triangle SIV and mdx cells additionally
- confirms the specificity of the anti-Na_v1.5 antibody since these findings accord with previously reported
- results^{5,7}. In addition, they show that the reduction in whole-cell I_{Na} in mdx cells is at least partly due to reduced
- Na_v1.5 expression at the lateral membrane⁸.
- To assess the effects of Na_v1.5 truncation and dystrophin deficiency on Na_v1.5 expression at the crest and the
- groove, we determined the distances from each α -actinin cluster to the closest Na_v1.5 cluster in mdx and Δ SIV
- lateral-membrane images, and compared them to wild-type values. The frequency histogram (**Figure 3e**) shows
- that the proportion of Na_v1.5 clusters within 50 nm of α -actinin is reduced by ~25% in *mdx* and Δ SIV compared
- to wild-type cells. When the proportion of $Na_v 1.5$ within 50 nm of α -actinin is plotted for each individual cell
- (Figure 3f), only Δ SIV cardiomyocytes show a significant ~20% reduction compared to wild type. Although
- mean $Na_v 1.5$ expression in mdx cells is reduced by about 17% compared to wild type, the data show high
- variability (**Figure 3f**), which correlates with the high variability in Na_v1.5 expression at the lateral membrane
- of mdx cells (**Figure 3d**). The affinity of Na_v1.5 for the groove is however similar in wild-type, mdx, and Δ SIV
- cells when comparing the proportion of $Na_v 1.5$ clusters within 50 nm of α -actinin from experimental images,
- and random and high-affinity simulations for each genotype (**Supplementary Figure 3a**).
- Taken together, these results establish that (1) $Na_v 1.5$ expression at the groove of the lateral membrane partially
- depends on the Δ SIV motif and dystrophin expression; (2) in dystrophin-deficient cells, Na_v1.5 clusters at the
- lateral membrane are organized into simpler shapes compared to wild-type cells; and (3) dystrophin deficiency
- induces high variability in Na_v1.5 expression and Na_v1.5 cluster size.

NA_V1.5 IS EXPRESSED IN THE T-TUBULAR MEMBRANE

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- 29 Na_v1.5 expression in the T-tubules was assessed by co-staining cardiomyocytes for Na_v1.5 and the T-tubular
- marker Bin1, and imaging an intracellular plane of each cell in highly inclined and laminated optical (HILO)

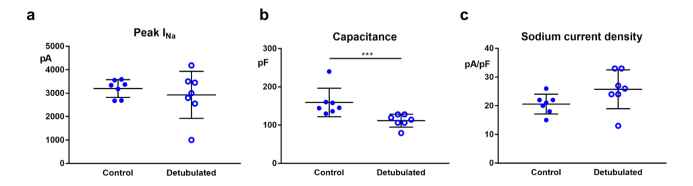


Figure 5. Whole-cell sodium current recordings in detubulated and normal cardiomyocytes. Compared to control cells, maximum sodium current (a) didn't change, capacitance (b) was reduced by about 30%, and sodium current density (c) increased by about 25% (non-significant) in detubulated cells. N = 5, n = 7 for both conditions. ***, p = 0.006, Mann-Whitney test.

mode with the TIRF objective. Bin1 was chosen as a T-tubular marker since it binds and shapes the T-tubular membrane and assists in the trafficking and clustering of the voltage-gated calcium channel $Ca_v 1.2^{24,26}$. The Bin1 antibody was first validated by confirming that Bin1 stainings are in close proximity to α -actinin, given that T-tubules run in close proximity to the sarcomeric Z-disc (N = 3, n = 59) (**Supplemental figure 4**)¹³. Then, we assessed whether $Na_v 1.5$ and Bin1 associated randomly or not by performing simulations for each SMLM image (N = 3, N = 39), where $Na_v 1.5$ clusters were redistributed either randomly or with high affinity for Bin1 (**Figure 4a-c**). $Na_v 1.5$ clusters at the lateral membrane and intercalated disc were excluded. Then, the distance from each Bin1 cluster to the closest $Na_v 1.5$ cluster was determined. The frequency histogram shows that about 16-17% of $Na_v 1.5$ clusters is within 50 nm of Bin1 in both original images and high-affinity simulations, whereas in random simulations this value was only ~8% (**Figure 4d**). This was confirmed when plotting the proportion of $Na_v 1.5$ clusters within 50 nm of Bin1 for each individual image (**Figure 4e**). Together, these findings indicate that a considerable subset of intracellular $Na_v 1.5$ is expressed in the T-tubular membrane.

T-TUBULAR SODIUM CURRENT CANNOT BE ASSESSED BY WHOLE-CELL RECORDINGS

Having shown that Na_v1.5 is expressed at the T-tubules (**Figure 4**), we investigated whether we could record a T-tubular sodium current. To this end, we compared whole-cell I_{Na} recordings from normal (N = 5, n = 7) and detubulated (N = 5, n = 7) ventricular cardiomyocytes. Firstly, we concluded that detubulation was successful as the capacitance of detubulated cells was 30% lower than of control cells (**Figure 5b**). Whole-cell I_{Na} did not decrease after detubulation (**Figure 5a**), which indicates that the vast majority of sodium current is conducted by sodium channels at the intercalated disc and lateral membrane. Comparing the current density shows a 25% yet non-significant increase in I_{Na} density after detubulation (**Figure 5c**), further confirming that the majority of sodium channels is outside the T-tubular domain. As discussed in the introduction, current densities of

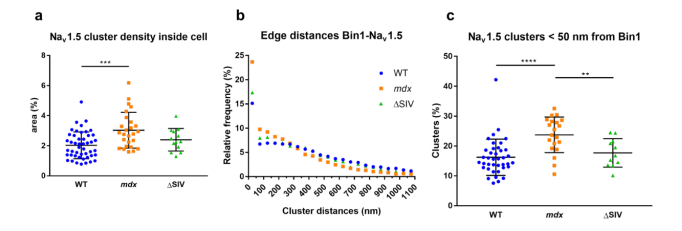


Figure 6. Na_v**1.5 expression is increased in T-tubules of** *mdx* **mice. a**, Na_v**1.5** cluster density in intracellular membrane compartments and T-tubules is ~30% higher in *mdx* cardiomyocytes than in wild-type cardiomyocytes, but no change was observed in ΔSIV compared to wild-type cells. **b**, For each Bin1 cluster, the distance to the closest Na_v**1.5** cluster is measured. Cluster distances are plotted in a frequency histogram with bins of 50 nm. The given histogram indicates that Na_v**1.5** preferentially locates within 50 nm to Bin1, in ΔSIV and *mdx* more so than in wild-type cells. **c**, In *mdx* cells, a higher proportion of Na_v**1.5** clusters is within 50 nm from Bin1 than in wild-type cells. Together with the increase in overall intracellular Na_v**1.5** expression in *mdx*, this indicates that T-tubules in *mdx* cells show a higher Na_v**1.5** expression. ΔSIV cells show a similar T-tubular Na_v**1.5** expression as WT cells. **a**, Wild type *N* = 3, n = 47; *mdx* N = 3, n = 26; ΔSIV N = 3, n = 13. **b**,**c** Wild type N = 3, n = 39; *mdx* N = 3, n = 20; ΔSIV N = 3, n = 11. **, p = 0.0095; ****, p = 0.0002; *****, p < 0.0001, Mann Whitney test.

- 1 detubulated and control cardiomyocytes cannot be compared directly as capacitance measurements in control
- 2 cardiomyocytes result in a considerable underestimation of cell membrane area. Thus, our data show that the T-
- tubular I_{Na} is below the detection limit of the whole-cell patch-clamp technique.

NA_V1.5 EXPRESSION IS INCREASED IN T-TUBULES FROM DYSTROPHIN-DEFICIENT CARDIOMYOCYTES, AND

CLUSTER ORGANIZATION DEPENDS ON THE SIV MOTIF AND DYSTROPHIN

- We next investigated whether dystrophin deficiency (mdx) and deletion of the SIV motif of Na_v1.5 affected
- Na_v1.5 expression in the T-tubules. Firstly, we determined that overall Na_v1.5 cluster density in intracellular
 - planes was increased in mdx compared to wild-type cells, whereas no difference was observed between ΔSIV
- and wild-type cells (**Figure 6a**; wild type N = 3, n = 47; mdx N = 3, n = 26; $\Delta SIV N = 3$, n = 13). We also noted
 - an increase in cluster solidity in mdx and an increase in cluster circularity in mdx and ΔSIV compared to wild-
 - type cells, while cell size and average Na_v1.5 cluster size did not differ between the three genotypes. Similar to
 - what we observed at the lateral membrane, this indicates that intracellular Na_v1.5 clusters have geometrically
- simpler shapes in mdx and ΔSIV cells.

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- The frequency histogram plotting distances between Bin 1 and $Na_v 1.5$ shows that the proportion of $Na_v 1.5$ within
- 50 nm to Bin1 was \sim 30% higher in mdx and \sim 4% higher in Δ SIV compared to wild type (**Figure 6b**; wild type
- N = 3, n = 39; mdx N = 3, n = 20; ΔSIV N = 3, n = 11). Plotting these values from individual cells confirmed
- this increase of T-tubular Na_v1.5 in mdx cells, but no difference between Δ SIV and wild type cells was observed

- 1 (**Figure 6c**). When comparing T-tubular Na_v1.5 expression from original images and random and high-affinity
- simulations, Na_v1.5 displays an affinity for Bin1 in all three genotypes; but in *mdx* cells, Na_v1.5 has an even
- 3 higher affinity for Bin1 than in the high-affinity simulations (**Supplementary figure 3b**).
- 4 Taken together, these data indicate that dystrophin deficiency induces a higher Na_v1.5 expression associated
- 5 with intracellular membrane compartments and T-tubules compared to wild type, whereas this effect was not
- observed in Δ SIV cells. Intracellular Na_v1.5 cluster shape were also markedly simpler in Δ SIV and mdx
- 7 compared to wild-type cardiomyocytes.

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8 DISCUSSION

This work aimed to surpass the previously published limited-resolution characterization of Na_v1.5 expression 9 and cluster organization at the lateral membrane and T-tubules of cardiomyocytes. To this end, we applied novel 10 quantitative single-molecule localization microscopy (SMLM) and modeling techniques to investigate Nav1.5 11 organization at the crest, groove, and T-tubules in cardiomyocytes from wild-type mice, dystrophin-deficient 12 (mdx) mice, and mice expressing C-terminally truncated Na_v1.5 (Δ SIV). We showed that (1) Na_v1.5 expression 13 in the groove of the lateral membrane partly depends on dystrophin and the SIV motif of Na_v1.5; (2) dystrophin 14 is involved in Na_v1.5 cluster organization at the lateral membrane; (3) Na_v1.5 is expressed in the T-tubules, 15 although we could not assess T-tubular sodium current with our electrophysiological approach; (4) T-tubular 16 Na_v1.5 expression is increased in dystrophin-deficient but not in ΔSIV cardiomyocytes; and (5) intracellular 17 Na_v1.5 cluster organization depends on dystrophin and the SIV motif. These findings are schematically 18 19 summarized in **Figure 7**.

We convincingly showed that $Na_v1.5$ is expressed in the T-tubules by co-staining $Na_v1.5$ with the T-tubular marker Bin1 in SMLM recordings, and assessing the robustness of this association by novel *in silico* methods^{25,27}. By comparing original images to simulations of these images in which $Na_v1.5$ clusters were redistributed either randomly or with high affinity for Bin1, the T-tubular population of $Na_v1.5$ in original images is twice as high as in random populations and comparable to that from high-affinity simulations (**Figure 4e**). The function of $Na_v1.5$ at the T-tubules remains elusive, although $Na_v1.5$ in the T-tubules will theoretically increase conduction velocity²⁸. Since some Bin1 and $Na_v1.5$ clusters overlapped (**Figure 4a**), we may hypothesize that Bin1 could be involved in the regulation of $Na_v1.5$ as Bin1 also regulates the trafficking of voltage-gated calcium channel $Ca_v1.2^{29}$.

We showed that the presence of a T-tubular I_{Na} could not be deducted from whole-cell electrophysiological methods: whole-cell currents between detubulated and control cardiomyocytes were similar (**Figure 5a**). The

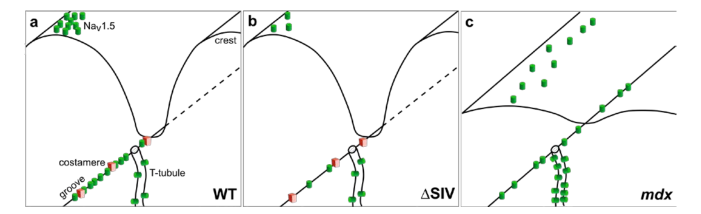


Figure 7. Model depicting changes at the lateral membrane of ΔSIV and mdx cardiomyocytes. Compared to the wild-type condition (a), ΔSIV cells (b) show less Na_v1.5 at the lateral membrane overall and specifically in the groove, whereas the lateral membrane structure remain intact. In mdx cells (c), the lateral membrane is flatter than in wild-type and ΔSIV cells, and Na_v1.5 expression at the lateral membrane is reduced overall; yet the expression at groove of mdx cells is not as affected as that of ΔSIV cells. Costameres are severely compromised in mdx cells. In the T-tubules, Na_v1.5 is similarly expressed in wild-type (a) and ΔSIV cells (b), and expression is elevated in mdx cardiomyocytes (c).

T-tubular I_{Na} may not surpass the noise of whole-cell recordings, or the T-tubular membrane may not be properly voltage clamped in our whole-cell approach, thus a proportion of T-tubular sodium channels may not be activated (space clamp issue). In the past, the T-tubular I_{Na} was suggested to be ~30% of total I_{Na} by comparing current densities from detubulated and control cells¹⁶. Measuring the capacitance in control cells underestimates cell membrane area, however, which disqualifies the direct comparison of current densities non-detubulated to detubulated cells¹⁷. Other electrophysiological methods have not been able to record T-tubular currents either: cell-attached²⁰ and SICM data^{11,30} only show currents from a few voltage-gated calcium channels whereas a much stronger calcium current would be expected from the T-tubules. To assess the T-tubular I_{Na} , more specific methods need to be developed.

When dystrophin is absent, less $Na_v1.5$ is expressed at the lateral membrane (**Figure 3a**), but more $Na_v1.5$ is expressed in the T-tubules compared to wild-type cells (**Figure 6c**). The reduction $Na_v1.5$ at the lateral membrane of mdx cells may be explained by the loss of costamere integrity (**Figure 7**)³¹. We may also hypothesize that dystrophin deficiency causes a partial rerouting of $Na_v1.5$ trafficking to the T-tubules instead of to the lateral membrane by an unknown mechanism. The short dystrophin product Dp71 may be involved since it is present only at the T-tubules³², and binds ion channels and the cytoskeleton³³. In light of the overall reduction of Nav1.5 protein expression and I_{Na} in mdx hearts⁸, the relative increase of T-tubular $Na_v1.5$ expression may make matters worse: due to the restricted space in the T-tubular lumen, the T-tubular sodium current may partially self-attenuate^{6,34}. A *de facto* translocation of $Na_v1.5$ from the lateral membrane to the T-tubules of mdx cells may quench whole-cell sodium current and contribute to the reduction in ventricular conduction velocity observed in mdx mice^{6,8}. Whether this effect indeed takes place *in vivo* requires further research.

1 Interestingly, $Na_v 1.5$ cluster shapes in the groove and the intracellular compartment are simpler in mdx than in

wild-type cells, illustrated by the increased solidity and circularity (**Supplementary Figure 2**). This indicates

that dystrophin is involved in scaffolding and shaping the Na_v1.5 clusters, which may be related to the large size

(427 kD) of dystrophin, and to the link to the actin cytoskeleton dystrophin provides³⁵. On the other hand, Na_v1.5

clusters in Δ SIV cells show an increase in solidity only in intracellular planes (**Supplementary Figure 2b**). This

suggests that the interaction of Na_v1.5 with the syntrophin-dystrophin plays a role in cluster organization in the

intracellular compartment/T-tubules, while at the lateral membrane, the secondary N-terminal syntrophin-

binding site of Na_v1.5 may support interaction with dystrophin and wild-type-like Na_v1.5 cluster organization³⁶.

The residual $Na_v 1.5$ expression at the lateral membrane of mdx mice may be explained by the dystrophin

homologue utrophin³⁷. Utrophin is expressed during the fetal phase, and re-expressed in adult tissue when

dystrophin is absent in mice, but not in humans^{37,38}. Since we found that Na_v1.5 cluster density and size at the

lateral membrane and at the groove is highly variable in *mdx* cells (**Figure 3a,c, Supplementary Figure 2a**),

ventricular cardiomyocytes may display a high variability in compensatory utrophin and Dp71 expression.

Utrophin however does not rescue a wild-type-like phenotype: the costameres and sarcolemma of utrophin-

expressing mdx mice are still weak³⁹, and the crest-groove profile at the lateral membrane of mdx cardiomyocytes

is flattened compared to wild type (**Figure 7c**)⁴⁰. The effects of dystrophin deficiency when the aforementioned

compensatory mechanisms are not available will need to be assessed in samples from the hearts of Duchenne

patients and of dystrophin/utrophin/Dp71 triple knock-out mice.

Like mdx mice, ΔSIV mice express less Na_v1.5 at the lateral membrane, and the groove pool is specifically

affected (**Figure 3d,f**). The Δ SIV truncation reduces the interaction of Na_v1.5 with the syntrophin-dystrophin

complex⁵. Since ΔSIV mice still express dystrophin, costameres presumably remain intact. Of note, the

syntrophin-dystrophin complex may also interact with the N-terminus of Na_v1.5³⁶, possibly explaining the

reduction, rather than the abolition, of Na_v1.5 localization to the groove and the lateral membrane. Contrary to

mdx mice, we did not observe an increase of Na_v1.5 at the T-tubules in Δ SIV mice, which may be explained by

their intact costameres.

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A groove-specific reduction of $Na_v1.5$, as shown for ΔSIV and a subset of mdx cardiomyocytes, and an increase

in T-tubular $Na_v 1.5$, as shown for mdx cells, may affect conduction and excitability in different ways. However,

late openings between the crest and the groove as well as biophysical properties did not differ in murine

cardiomyocytes¹¹. Future studies should address specific differences in function and regulation of Na_v1.5 in the

crest, groove, and T-tubules.

Intriguingly, while our data revealed that Na_v1.5 is reduced at the groove of mdx and Δ SIV mice, Rivaud *et al.*

have recently shown in a murine heart failure model (transient aortic constriction) that sodium current is

specifically reduced at the crest, not at the groove, compared to sham-operated animals¹¹. Rivaud *et al.* showed

that Na_v1.5 cluster size, not cluster density, was markedly reduced at the lateral membrane in the heart failure

model, whereas our data show that cluster density, not cluster size, was reduced in mdx and ΔSIV cells

(Supplementary Figure 2a). Expression and organization of Na_v1.5 at the crest and the groove was however

not investigated in this heart failure model. Together, these data suggest that cardiomyocytes respond differently

to dystrophin deficiency, Na_v1.5 truncation (ΔSIV), and transient aortic constriction¹¹. One obvious difference

is the hypertrophy observed in the heart failure model but not in mdx or ΔSIV mice. These different pathways

remain to be elucidated in detail.

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8 The phenotype of Brugada syndrome patients carrying the mutation p.V2016M, which changes the C-terminal

9 motif SIV into SIM, may be partly explained by a reduction in Na_v1.5 cluster density in the groove of the lateral

membrane and a reduction of dystrophin-Na_v1.5 interaction⁸. Indeed, in Δ SIV cardiomyocytes, transversal

conduction velocity, total sodium current, and sodium current at the lateral membrane are decreased⁵, which

may contribute to the Brugada syndrome phenotype. However, the specific role of groove-specific Na_v1.5

reduction remains to be determined.

A limitation of our study is that mdx mice display various compensatory mechanisms discussed above that do

not allow us to assess the effect of a "pure" dystrophin deficiency. Mdx mice moreover do not have the optimal

controls, since they were on a Bl6/Ros background while being backcrossed to Bl/6J in the fourth generation,

whereas wild-type and Δ SIV mice were littermates on a pure Bl6/J background. The discrepancy in backgrounds

may have introduced variability in our data, although the data are consistent with previously published results^{7,8}.

Lastly, our methods did not allow us to quantify crest expression of Na_v1.5.

Taken together, our findings provide important and previously unattainable mechanistic insights on the

properties of Na_v1.5 organization on the crest, groove, and lateral membrane of cardiomyocytes, and are an

important step towards identifying cardiac domain-specific molecular determinants of Na_v1.5 and location-

specific effects of SCN5A mutations. However, pool-specific function and regulation of Na_v1.5 at the crest,

groove, and T-tubules remains to be studied in more detail.

25 **METHODS**

ETHICAL STATEMENT

- 27 All animal experiments conformed to the Guide to the Care and Use of Laboratory Animals (US National Institutes of
- Health, publication No. 85-23, revised 1996); were approved by the Cantonal Veterinary Administration, Bern,
- 29 Switzerland; conformed to the New York University guidelines for animal use and care (IACUC Protocol 160726-03 to
- 30 MD, approved 07/11/2018); and have complied with the Swiss Federal Animal Protection Law.

MOUSE MODELS

1 Dystrophin knock-out (mdx^{5CV}) mice

- The mdx^{5CV} mouse strain displays total deletion of dystrophin protein. It was generated as described previously⁴¹, and
- 3 purchased from the Jackson laboratory (Bar Harbor, Maine; stock #002379). Mdx mice were on a mixed background of
- 4 Bl6/Ros and Bl6/J (fourth generation of backcrossing). Male littermates aged 16 weeks were used.

5 **ΔSIV knock-in mice**

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- 6 In ΔSIV knock-in mice, the C-terminal SIV motif of Na_v1.5 is deleted (*Scn5a*-p.S2017STOP), as described previously⁵.
- 7 ΔSIV mice and wild-type littermates are on a pure Bl6/J background. All mice (male and female) were 16 weeks old.

ISOLATION OF MURINE VENTRICULAR MYOCYTES

- 9 All experimental steps are performed at room temperature unless specified otherwise.
- 10 Cardiomyocytes were isolated based on a previously established enzymatic method⁸. Briefly, mice were euthanized by
- 11 cervical dislocation. Hearts was excised, cannulated, and mounted on a Langendorff system for retrograde perfusion at
- 12 37°C. Hearts were rinsed first with nominally Ca²⁺-free solution containing (in mM) 135 NaCl, 4 KCl, 1.2 MgCl₂, 1.2
- NaH₂PO₄, 10 HEPES, and 11 glucose (pH 7.4, NaOH adjusted), then with digestion solution for 15 minutes, consisting of
- the aforementioned solution supplemented with 50 μM Ca²⁺ and collagenase type II (1 mg/mL; Worthington, Allschwil,
- Switzerland). Subsequently, atria were removed. Ventricles were transferred to nominally Ca²⁺-free solution and minced
- into small pieces. To increase the yield of isolated cells, digested ventricular tissue was triturated gently and filtered through
- 17 a 100 μm nylon mesh. Before use, cells were subjected to a calcium increase procedure.

18 CONFOCAL MICROSCOPY

- 19 Isolated cardiomyocytes were fixed with acetone at -20°C for 10 minutes in cell culture chambers. Cells were washed twice
- with phosphate-buffered saline (PBS), permeabilized with 0.1% saponin in PBS for 2 x 7 minutes, and blocked for 15
- 21 minutes with PBS containing 10% normal goat serum, 1% bovine serum albumin (BSA), 0.1% saponin, and 30 μg/mL Fab
- fragment donkey anti-mouse IgG (H+L) (Jackson Immuno Research, Baltimore, Maryland). All following antibody
- dilutions and washing steps were performed with incubation buffer (3% normal goat serum, 0.05% saponin, and 1% BSA
- in PBS) unless specified otherwise. Primary antibodies were diluted and applied onto cells for overnight incubation at 4°C.
- 25 Afterwards, cells were washed 4 times for 5 minutes, and incubated with secondary antibodies for 2 hours. Cells were
- 26 washed 3 times for 5 minutes, incubated with DAPI (1 μL in 200 μL PBS) for 20 minutes, washed twice with PBS, and
- 27 finally embedded in FluorSave reagent (Merck, Burlington, Massachusetts). Cells were imaged on a confocal microscope
- 28 (LSM710, Zeiss, Oberkochen, Germany).

SINGLE-MOLECULE LOCALIZATION MICROSCOPY

Sample preparation

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- Freshly isolated cardiomyocytes were plated on laminin-coated glass coverslips and allowed to adhere at 37°C for 30
- minutes. Cells were fixed with 4% paraformaldehyde for 10 minutes and washed 3 times with PBS. To quench

- autofluorescence, cells were placed overnight under 470 nm LED light. Then, cells were permeabilized with 0.1% Triton
- 2 X-100 in PBS for 10 minutes, and blocked with blocking solution (2% glycine, 2% BSA, and 0.2% gelatin in PBS) for 30
- 3 minutes. All following antibody dilutions and washes were performed with blocking solution unless specified otherwise.
- 4 Unconjugated primary antibodies were applied to the cells for 1 hour, followed by three 5-minute washes. Secondary
- 5 antibodies were applied for 15 minutes, followed by three 5-minute washes and an optional 15-minute incubation with
- 6 conjugated primary antibodies. Lastly, cells were washed with PBS for 5 minutes. Coverslips were mounted onto
- 7 microscope slides in which two small holes were drilled and using double-sided tape as spacers, creating a fluid chamber.
- 8 Chambers were sealed with epoxy resin.

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Optical setup and image acquisition

- SMLM imaging was performed in accordance to a previously described method²³. To achieve stochastic fluorophore
- blinking, imaging buffer was added to the cells, containing 200 mM 2-mercaptoethylamine, 10% glucose, 0.04 mg/mL
- 12 glucose oxidase and 0.08 mg/mL catalase in T50 buffer (10 mM Tris pH 8 and 50 mM NaCl in MilliQ). Samples were
- imaged on a customized Leica DMI 300 inverse microscope equipped with an HCX PL APO 63X NA = 1.47 OIL CORR
- 14 TIRF objective (Zeiss), a 2X tube lens (Diagnostic Instruments, Sterling Heights, Michigan) and a chromatic aberration
- correction lens (AC254-300-A; Thorlabs, Newton, New Jersey). Samples were sequentially excited by a 639 nm laser (,
- MRL-FN-639-800; UltraLasers, Newmarket, Canada), 561 nm laser (MGL-FN-561-200; UltraLasers), and 488 nm laser
- 17 (OBIS; Coherent, Santa Clara, California). A 405 nm laser (MDL-III-405-150; CNI, Changchun, China) was used to
- 18 reactivate AlexaFluor 647 fluorophores. Lasers were aligned by a penta-edged dichroic beam splitter
- 19 (FF408/504/581/667/762-Di01-22x29). The 488, 561, and 639 laser lines were adjusted to ~0.8, 1.0, and 1.5 kW/cm²,
- 20 respectively. The emitted fluorescence was filtered by the applicable single-band fluorescence filter (FF01-531/40, FF01-
- 21 607/36, and FF01-676/37 for AlexaFluor 488, 568, and 647, respectively; Semrock, Rochester, New York) in a filter wheel,
- and recorded at 33 Hz by a sCMOS camera (Prime95B; Photometrics, Tucson, Arizona) with a minimum of 2000 frames
- per wavelength. Recordings were controlled by MicroManager software (version 1.4.22; www.micro-manager.com). The
- readout noise of each camera pixel was pre-calibrated and characterized by a Gaussian distribution⁴².

Alignment of different colors for SMLM images

- 26 Different colors were aligned following previously described methods²³. Briefly, broad-spectrum fluorescent beads
- 27 (diameter ~100 nm, TetraSpec; Thermo Fisher, Waltham, Massachusetts) were imaged sequentially in the blue (488 nm),
- green (568 nm), and red (639 nm) channels. The images from the blue and green channels were aligned to the red channel
- using a second polynomial warping algorithm in Matlab (version R2017b; Mathworks, Natick, Massachusetts).

Single-molecule localization

- 31 Single-molecule localization also followed a previously described method²³. In short, from each image stack, each frame
- was box-filtered. Box size was four times the full width at half maximum of a 2D Gaussian point spread function (PSF).
- 33 Each pixel was weighted by the inverse of its variation during such box-filtering. The low-pass filtered image was extracted
- 34 from the raw image, and local maxima were identified. Local maxima from all frames of one image stack were subjected
- to 2D Gaussian multi-PSF fitting⁴³. The 2D Gaussian single-PSF fitting was performed using the maximum likelihood
- estimation (MLE) algorithm on a GPU (GTX 1060, CUDA 8.0; Nvidia, Santa Clara, California). The likelihood function

- 1 at each pixel was based on convolving the Poisson distribution of the shot noise, which is governed by the photons emitted
- 2 from nearby fluorophores, and the Gaussian distribution of the readout noise characterized by the pre-calibrated
- 3 expectation, variation, and analog-to-digital conversion factor. Fitting accuracy was estimated by Cramér-Rao lower bound.

Image analysis

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- 5 Images were processed based on previously described methods²⁷. Briefly, images were subjected to a smoothing filter,
- 6 adjusted for brightness and contrast, and filtered to a threshold to obtain a binary image with ImageJ software (version
- 7 1.51j8; https://imagej.net). Regions of interest were drawn, defined as regions with regular α-actinin and/or Bin1 staining,
- 8 depending on the applied antibodies. We excluded the intercalated disc regions from both surface and intracellular
- 9 recordings, and the lateral membrane regions from intracellular recordings. Clusters were analyzed in ImageJ. Image
- simulations were performed using the Interaction Factor plugin in ImageJ²⁵, and distance between clusters in experimental
- and simulated images were obtained using the ImageJ function 'Analyze particles' and a script written in Python (version
- 12 2.7; www.python.org) that utilized the image processing packages 'scikit-image' and 'mahotas'²⁷. Frequency histograms
- of cluster distances were made in GraphPad Prism (version 7; GraphPad Software, San Diego, CA, USA). The investigator
- that analyzed the images was ignorant of the corresponding genotypes.

ANTIBODIES

- Primary antibodies were: custom-made rabbit anti-Na_v1.5 (epitope: amino acids 493-511 of rat Na_v1.5; Pineda Antibody
- Service, Berlin, Germany; 1:100), mouse anti-Bin1 (amphiphysin-II; epitope: amino acids 179-207 of human Bin1; Santa
- 18 Cruz Biotechnology, Dallas, Texas; 1:50), mouse anti-α-actinin (raised against rabbit skeletal α-actinin; Sigma-Aldrich, St.
- Louis, Missouri; 1:400), and the same anti-α-actinin antibody directly conjugated to AlexaFluor 488 or 647 (Thermo Fisher;
- 20 1:2000 or 1:5000, respectively). Secondary antibodies were: rabbit AlexaFluor 568 (Life Technologies, Carlsbad,
- California; SMLM: 1:10,000; confocal: 1:200), mouse AlexaFluor 647 (Life Technologies; SMLM: 1:5000), and mouse
- 22 AlexaFluor 488 (Life Technologies; confocal: 1:200).

23 **DETUBULATION**

- Isolated cardiomyocytes were detubulated by osmotic shock using formamide based on a previously described method⁴⁴.
- 25 Briefly, cardiomyocytes were resuspended in 2 mL 1.5 M formamide solution for 15 minutes and washed twice with
- 26 extracellular solution (see "Electrophysiology" section).

ELECTROPHYSIOLOGY

- Whole-cell sodium currents (I_{Na}) were recorded at 22-23°C using a VE-2 amplifier (Alembic Instruments Inc., Montréal,
- Canada). Borosilicate glass pipettes were pulled to a series resistance of $\sim 2 \text{ M}\Omega$. Recordings were visualized with pClamp
- software (version 8, Axon Instruments, Union City, California), and analyzed with pClamp and OriginPro (version 7.5,
- OriginLab Corp., Northampton, Massachusetts). Current densities (pA/pF) were calculated by dividing the peak current by
- the cell capacitance. Cells were bathed in extracellular solution containing (in mM) 5 NaCl, 125 NMDG-Cl, 5.6 CsCl, 5
- BaCl2, 1 MgCl2, 10 HEPES, and 5 glucose (pH 7.4, CsOH adjusted). Internal solution contained (in mM) 130 KCl, 4 Mg-

34 ATP, 12 NaCl, 1 MgCl₂, 1 CaCl₂, 10 EGTA, 10 HEPES (pH 7.2, KOH adjusted).

STATISTICS

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- Data are presented as means \pm standard deviation. Differences between two groups were analyzed in GraphPad Prism by
- 3 two-tailed T-tests if the data were normally distributed, and Mann-Whitney tests if the data were not. P < 0.05 was
- 4 considered statistically significant. No explicit power analysis was used. Three (SMLM) or five (electrophysiology) mice
- 5 per genotype were used as biological replicates, providing sufficient statistical power. For SMLM, 47/24 (wild type), 26/27
- 6 (mdx), and 13/19 (Δ SIV) cells were imaged on intracellular and surface imaging planes, respectively, also providing
- 7 sufficient statistical power. For individual experiments, cells from different genotypes were always isolated, or stained and
- 8 imaged side by side. We performed 5/4 (wild type), 3/3 (mdx), and 2/3 (ΔSIV) technical replicates for intracellular and
- 9 surface imaging, respectively. A technical replicate is defined as one round of staining and imaging. For electrophysiology,
- 10 currents were recorded in seven control and seven detubulated cells from five wild-type mice. Each mouse counts as one
- technical replicate as data from each mouse were collected on one experiment day.

Exclusion and inclusion criteria

- 13 Regarding SMLM data, cells with irregular Bin1 staining were excluded, as they indicated a technical problem and/or T-
- tubular remodeling. All other cells were included as α-actinin and Na_v1.5 stainings showed consistent results. No outliers
- 15 were excluded.
- Regarding electrophysiology data, we only included recordings with gigaseal and high-quality voltage clamp. Among those,
- 17 no outliers were excluded.

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AUTHOR CONTRIBUTIONS

- SHV, HA, and JSR contributed conception of the study; SHV, JSR, EAP, ER, and MD contributed methodology;
- 25 SHV and JSR performed experiments; SHV contributed validation, data curation, project administration,
- visualization, and writing-original draft; SHV and JSR performed formal analysis and investigation; EAP, ER,
- MD, and HA contributed resources; JSR, EAP, ER, MD, and HA supervised; SHV and HA contributed funding
- acquisition; and all authors contributed writing–review & editing.



- 2 **Competing interest:** The authors declare that the research was conducted in the absence of any commercial or
- 3 financial relationships that could be construed as a potential conflict of interest.

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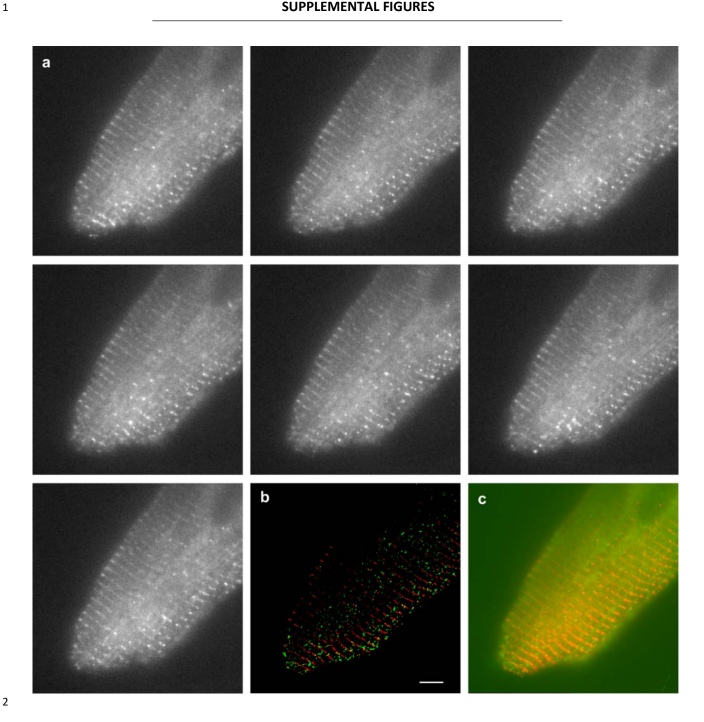
25

1 SOURCE DATA

- Figure 2d Source data 1. This spreadsheet contains the percentages of Na_v1.5 clusters which edges are 50
- nm from the edge of the nearest α -actinin cluster at the lateral membrane of wild-type cardiomyocytes, and the
- 4 corresponding statistical analyses. Values from original images are compared to those from simulations in which
- 5 Na_v1.5 clusters are redistributed over the respective image either randomly or with high affinity for α-actinin.
- 6 Figure 3d-f Source data 2. This spreadsheet contains data on Na_v1.5 cell surface expression (pertaining to
- 7 Fig 3d), frequency distribution of edge distances from any α-actinin cluster to the closest Na_v1.5 cluster
- 8 (pertaining to Fig 3e), and the percentage of Na_v1.5 clusters within 50 nm from α-actinin at the lateral
- 9 membranes of wild-type, mdx, and Δ SIV cells. Statistical analyses and descriptive statistics are given where
- 10 applicable.
- 11 **Figure 4d-e Source data 3.** This spreadsheet contains the values of the frequency distribution of edge
- distances from any Bin1 cluster to the closest Na_v1.5 cluster (pertaining to **Fig 4d**), and the percentages of Na_v1.5
- clusters within 50 nm from Bin1 (pertaining to **Fig 4e**) in intracellular recordings of wild-type cardiomyocytes.
- Statistical analyses and descriptive statistics are also given.
- Figure 5 Source data 4. This spreadsheet contains values of cell capacitance, maximum sodium current, and
- sodium current density recorded in normal and detubulated wild-type cardiomyocytes. Statistical analyses are
- included.
- Figure 6 Source data 5. This spreadsheet contains values of Na_v1.5 cluster density and frequency distributions
- of edge distances from any Bin1 cluster to the closest $Na_v1.5$ cluster in intracellular planes in wild-type, mdx,
- and Δ SIV cells. Statistical analyses and descriptive statistics are included.
- Supplemental figure 2 Source file 6. This spreadsheet contains values of cell size, average Na_v1.5 cluster
- size, $Na_v 1.5$ cluster solidity and circularity of wild-type, mdx, and ΔSIV cardiomyocytes at surface (pertaining
- to Supplemental Fig 2a) and intracellular imaging planes (pertaining to Supplemental Fig 2b). Statistical
- analyses are included.
- 25 **Supplemental figure 3 Source file 7.** This spreadsheet contains values of Na_v1.5 clusters within 50 nm from
- 26 α-actinin at the cell surface (pertaining to **Supplemental Fig 3a**) and Bin1 inside the cell (pertaining to
- 27 **Supplemental Fig 3b**). Values are compared between original images and simulations in which Na_v1.5 clusters
- are redistributed either randomly or with high affinity for α -actinin or Bin1, respectively. Statistical analyses are
- 29 also given.

30

SUPPLEMENTAL FIGURES



Supplemental figure 1. Fluorophores blink during SMLM image acquisition. a, Frames from a recording of a cardiomyocyte stained with anti-α-actinin antibodies directly conjugated to AlexaFluor 647 fluorophores and excited with a 639 nm laser. The fluorophores release photons intermittently - "blink" - due to the oxygen scavenging imaging buffer (see Methods). b, Reconstructed image depicting α -actinin in red and Na_v1.5 in green. Note that the fluorophores on the left side of the cell that did not blink, characterized by a faint striated pattern in all panels of (a), are not reconstructed in the final SMLM image (b). Scale bar 5 μ m.

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7

8

a Surface images

0.85

0.80

0.75

0.70

1

2

3

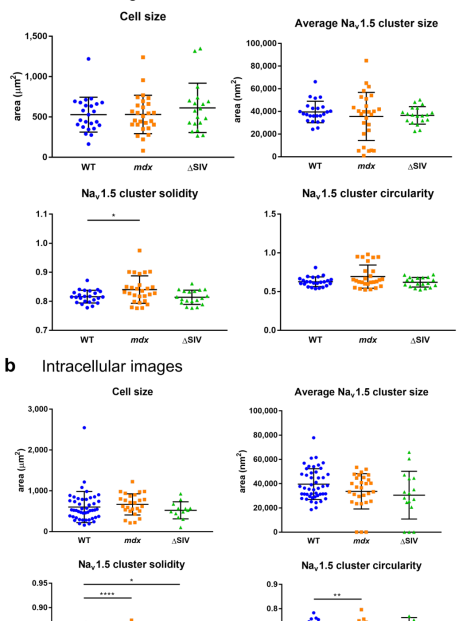
4

5

6

mdx

∆SIV



Supplemental figure 2. Cell and Na_v1.5 cluster properties of SMLM images. For all images from wild-type, mdx, and Δ SIV cells at the lateral membrane (a) and inside the cell (b), this figure shows cell size, and cluster size, circularity, and solidity of Na_v1.5 clusters per cell. Perfectly circular and solid clusters give value 1. Solidity indicates the ratio of the particle area to the area of the convex hull of the particle. a, Wild type N = 3, n = 24; mdx N = 3, n = 27; Δ SIV N = 3, n = 19. b, Wild type N = 3, n = 39; mdx N = 3, n = 20; Δ SIV N = 3, n = 11. *, p = 0.041 (a, solidity); p = 0.021 (b, solidity); **, p = 0.001; ****, p < 0.0001, Mann Whitney test.

WΤ

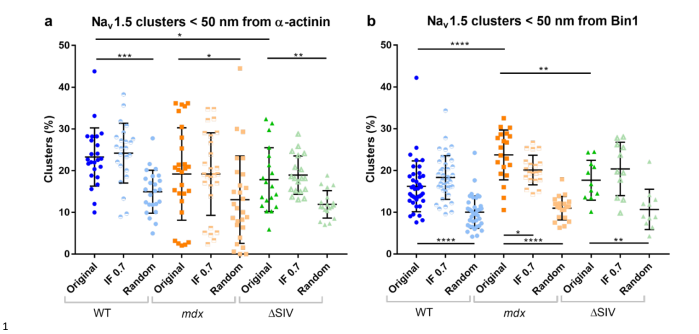
mdx

∆SIV

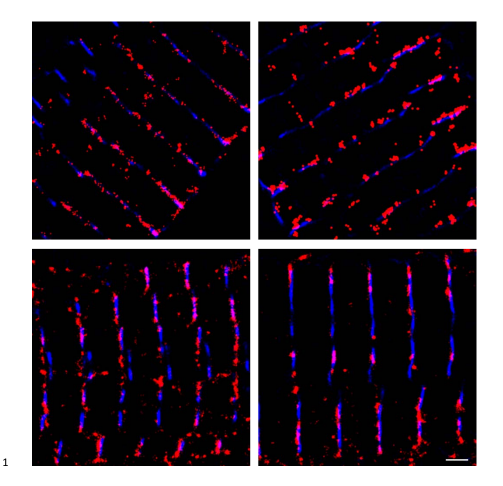
0.7

0.6

0.5



Supplemental figure 3. Comparing cluster distances of experimental SMLM images to simulations. Proportion of Na_v1.5 clusters within 50 nm from α -actinin at the lateral membrane (a) and Bin1 inside the cell (b). SMLM images from wild-type (blue), mdx (orange), and Δ SIV (green) cells are compared to their respective high affinity (IF 0.7) and random simulations. a, Wild type N = 3, n = 24; mdx N = 3, n = 27; Δ SIV N = 3, n = 19. b, Wild type N = 3, n = 39; mdx N = 3, n = 20; Δ SIV N = 3, n = 11. *, p = 0.0194 (a, Δ SIV-WT), p = 0.017 (b, mdx); ***, p = 0.0074 (a, Δ SIV) p = 0.0019 (b, Δ SIV), p = 0.0095 (b, mdx- Δ SIV); ***, p < 0.001, ****, p < 0.0001, unpaired T-test (b, wild type) and Mann Whitney test (all others).



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Supplemental figure 4. Bin1 antibody validation. Representative SMLM images of Bin1 (red) and α -actinin (blue) showing close proximity of Bin1 to α -actinin. This indicates that the anti-Bin1 antibody is most likely specific, as Bin1 binds the T-tubular membrane, and most T-tubules run along α -actinin lines, which constitute the sarcomeric Z-discs. Bin1 signal in between α -actinin lines may indicate axial T-tubular branches. N = 3, n = 57; scale bar 1 μ m.