

Pain Focus

Fluorescence labeling strategies for cell surface expression of TRPV1

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Regulation of ion channel expression on the plasma membrane is a major determinant of neuronal excitability, and identifying the underlying mechanisms of this expression is critical to our understanding of neurons. Here, we present two orthogonal strategies to label extracellular sites of the ion channel TRPV1 that minimally perturb its function. We use the amber codon suppression technique to introduce a non-canonical amino acid (ncAA) with tetrazine click chemistry, compatible with a transcyclooctene coupled fluorescent dye. Additionally, by inserting the circularly permutated HaloTag (cpHaloTag) in an extracellular loop of TRPV1, we can incorporate a fluorescent dye of our choosing. Optimization of ncAA insertion sites was accomplished by screening residue positions between the S1 and S2 transmembrane domains with elevated missense variants in the human population. We identified T468 as a rapid labeling site (~5 min) based on functional and biochemical assays in HEK293T/17 cells. Through adapting linker lengths and backbone placement of cpHaloTag on the extracellular side of TRPV1, we generated a fully functional channel construct, TRPV1exCellHalo, with intact wild-type gating properties. We used TRPV1exCellHalo in a single molecule experiment to track TRPV1 on the cell surface and validate studies that show decreased mobility of the channel upon activation. The application of these extracellular label TRPV1 (exCellTRPV1) constructs to track surface localization of the channel will shed significant light on the mechanisms regulating its expression and provide a general scheme to introduce similar modifications to other cell surface receptors.

Introduction

The non-selective cation channel TRPV1 is expressed in the small-diameter class of dorsal root ganglia neurons known as nociceptive c-fibers (Bevan and Szolcsányi, 1990; Caterina et al., 1997). Activation of TRPV1 in these nociceptors by temperatures above 43°C is linked to physiological heat sensation, and both oxidation and acidity acting on TRPV1 lead to the perception of noxious stimuli (Davis et al., 2000; Leffler et al., 2006; Chuang and Lin, 2009). Recent studies offer compelling evidence that TRPV1 function is central to inflammatory pain and, as such, has become a pharmacological target for pain intervention (Szolcsányi and Sándor, 2012).

As an important receptor for pain studies, TRPV1 channel research has made extensive progress along several aspects of its regulation, including desensitization by Ca²⁺, posttranslational modifications, lipid regulation, and trafficking (Koplas et al., 1997; Olah et al., 2001; Morenilla-Palao et al., 2004; Zhang et al., 2005; Vetter et al., 2008). This is neither a complete list of TRPV1 mechanisms of regulation nor are the listed attributes

necessarily mutually exclusive. In fact, the lipid PI(3,4,5)P3 is implicated as an essential signal for increased surface trafficking of TRPV1 (Stein et al., 2006; Stratiievska et al., 2018). Compelling evidence that surface expression of TRPV1 is an important aspect of the channel's regulation has motivated studies that monitor the trafficking of the channel (Morenilla-Palao et al., 2004; Zhang et al., 2005). Although voltage-clamp electrophysiology directly measures currents elicited from the total number of TRPV1 on the cell surface, there is an inherent ambiguity between current changes stemming from differences in singlechannel conductance versus differences in the total number of channels. As a complementary tool to electrophysiology, fluorescence imaging of cell surface channels and receptors fused with genetically encoded fluorescent proteins (FPs) is a bona fide approach to directly measure their changing numbers (Michaluk and Rusakov, 2022). By fusing a genetically encoded FP to the C-terminus of TRPV1, researchers can track the subcellular localization of the channel. Studies of TRPV1-FP surface

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expression in F11 DRG hybridoma cells and HEK293T/17 cells have informed our understanding about NGF-dependent trafficking and Ca²⁺-dependent surface mobility of TRPV1 (Stein et al., 2006; Senning and Gordon, 2015).

Two critical obstacles stand in the way of obtaining a more detailed mechanism behind TRPV1 trafficking and surface expression dynamics. The primary concern is that signals from fluorescently labeled channels on the surface must be discriminated from intracellular pools of the channel. If both populations of the channel are labeled with the same fluorescent protein, the signal-to-noise in the experiment is considerably less than if only the surface population were optically active. A secondary concern is that the fluorescent protein introduces a perturbation to the channel structure that may restrict aspects of its regulation (Wang et al., 2019). Moreover, channel function and interaction surfaces may be impacted by the inclusion of a large fluorescent protein domain.

Here, we present a comprehensive set of fluorescence research tools that are intended to address the limitations of current TRPV1 labeling strategies, especially as concerns trafficking and subcellular localization. Our fluorescent labeling strategies rely on a post-expression cell surface clickchemistry approach with cell impermeable fluorescent chemical ligands (Fig. 1 A). We have optimized the expression of a TRPV1-"tag" construct with an amber stop codon (tag) to introduce an amino acid-sized click-chemistry site with amber codon suppression techniques (Fig. 1, B i) (Zagotta et al., 2016; Blizzard et al., 2018; Bessa-Neto et al., 2021). Optimal placement of our tag site in the TRPV1 sequence was determined by genomic variant scanning, which builds on our previous work to relate variant positions identified in gnomAD with TRPV1 function (Karczewski et al., 2020; Mott et al., 2023). An alternative approach to extracellular labeling of TRPV1 was explored with the HaloTag click-chemistry system. We incorporated the circularly permuted HaloTag within an extracellular loop of TRPV1 (Fig. 1, B ii) (Deo et al., 2021). Although the TRPV1exCellHalo construct sustains a large increase in molecular mass, we show that the channel expresses well, functions similarly to wild-type TRPV1, and assembles into heteromeric channels with C-terminally labeled TRPV1 subunits.

Materials and methods

Bioinformatic analysis of human genomic data

To examine the TRPV1 sequence for variant amino acid positions, we accessed human genomic variant data from gnomAD 2.1.1. Human variant data for TRPV1 was downloaded directly from gnomAD servers (https://gnomad.broadinstitute.org; "export variants to CSV" option). Spreadsheet data were formatted in Excel (Microsoft) to read in the variant data to a Matlab (Mathworks) script, as done by Mott and coworkers, that generated a frequency plot for the number of missense variants at each residue position between TRPV1 positions 427 and 680 (see Fig. 2 A) (Mott et al., 2023). Synonymous mutations are ignored, and each occurrence of a missense variant is recorded as a variant in our counts. Residues with >100 variants were truncated

to 100 counts, and positional frequency was also smoothed with a boxcar average of three residues. Alignment with secondary structure was done with respect to rat TRPV1 residue positions, and the α -helical structure was based on the TRPV1 secondary structure assignment in Nadezhdin et al. (2021). Sequence alignment data for conservation analysis was obtained from Uniprot.org (https://www.uniprot.org), and alignments are displayed in Fig. 2 D with an alignment annotator (https://bioinformatics.org/strap/aa/).

Electrophysiology

Currents were recorded at room temperature in symmetric divalent-free solutions (130 mM NaCl, 3 mM HEPES, and 0.2 mM EDTA, pH 7.2) using fire-polished borosilicate glass pipettes with a filament (outer diameter 1.5 mm and inner diameter 0.86 mm; Sutter). Pipettes were heat-polished to a resistance of 3.5-7 $M\Omega$ using a micro forge. Cells were transfected with channel constructs and replated after 18-24 h on 12-mm coverslips and placed in a divalent-free solution in the chamber. 1 and 5 μM capsaicin solutions were prepared from a 5.3 mM capsaicin stock in ethanol. The same stock was diluted in the recording solution to 0.5 mM to make 0.03, 0.1, and 0.3 µM capsaicin solutions. Excised inside-out patches were positioned at the opening of a tube in a "sewer pipe" configuration with eight tubes connected to a rod controlled by an RSC-200 rapid solution changer (BioLogic). Capsaicin solutions (0.03, 0.1, 0.3, 1, and 5 μ M) were perfused for 20-30 s each as continuous 1-s sweeps stepped the voltage from resting potential (10 ms) to -80 mV (100 ms), then up to +80 mV (300 ms), and back to resting potential (10 ms). Currents were filtered at 5 kHz and recorded using an Axopatch 200A amplifier (Axon Instruments, Inc.) and PatchMaster software (HEKA). Data were analyzed in PatchMaster and IGOR (Wavemetrics). Individual patch-clamp recordings with a complete set of stable current measurements in all capsaicin solutions were normalized to the maximal current with 5 µM capsaicin. Normalized data were fitted to a Hill equation to arrive at the K_{1/2} parameter: $I = I_{\text{max}} \left[[\text{ligand}]^n / \left(K_{\frac{1}{2}}^n + [\text{ligand}]^n \right) \right]$, where n was permitted to float. Fig. 4 C reports the mean $K_{1/2}$ for patch doseresponse measurements made with TRPV1 wild-type and TRPV1exCellHalo. Statistical significance was determined by two-tailed Student's t test.

Cell culture

HEK293T/17 (CRL-11268; ATCC) cells were incubated in Dulbecco's modified Eagle medium, supplemented with 10% fetal bovine serum, 50 μg streptomycin, and 50 U/ml penicillin, at 37°C and 5% CO₂. Cells were passaged onto poly-L-lysine treated 25-mm coverslips. Cells were allowed at least 2 h to settle onto the slips before being transfected using Lipofectamine 2000 (Life Technologies) as described in the manufacturer's instructions. Cells transfected with TRPV1-tag mutants contained the target protein, TRPV1-tag-GFP in pcDNA3.1, an amber codon suppression plasmid, either pNEU (ratio 1:1), pHMEF5-H1U6-RS (ratio 1:3), or pAcBacl-NES-R284 (ratio 2:1), and the media was supplemented with either 10 μM LysZ (for pNEU), TCO (for pNEU or pHMEF5-H1U6-RS), or 30 μM Tetrazine3-Butyl (Tet3-Butyl for PAcBacl-



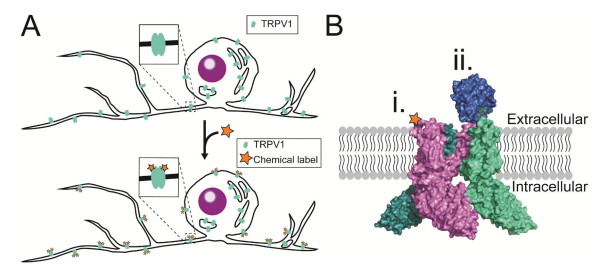


Figure 1. **Two strategies for extracellular labeling of TRPV1.** (A) TRPV1 expression in DRG neurons is distributed across the plasma membrane and intracellular compartments (top). If the extracellular side of TRPV1 can be labeled at a click-chemistry site, labeling the plasma membrane population of TRPV1 is possible (bottom). (B, i and ii) Amber codon suppression techniques permit the substitution of a ncAA at a single residue in the TRPV1 sequence. The addition of an ncAA with click-chemistry properties in an extracellular loop of TRPV1 allows surface labeling of the ion channel (PDB accession no. 7LQY; B, i). By introducing the HaloTag domain (blue; PDB accession no. 6U32) to an extracellular loop of TRPV1, cell-impermeable HaloLigands can be specifically targeted to the population of channels that are expressed on the plasma membrane (B, ii).

NES-R284) (Kita et al., 2016; Serfling et al., 2019; Jang et al., 2020). Following transfection, Ca²⁺ imaging experiments were carried out after 24 h, and Western blot experiments were completed 48 h after transfection.

Molecular biology

The TRPV1-exCellHalo construct was built using the standard Gibson Assembly method of cloning (NEB). Briefly, two primer sets were made for cpHaloTag in pCAG-HASAP (plasmid #138325; AddGene) and rTRPV1: fragment set 5'-AGGATGGGA AGAATAACTCTGCCTTTGCCCGCG-3'; 5'-GGTGTGGACTCCATA GGCAGCCCGGCAAATTCTGGCC-3' and vector set 5'-AATGGC CAGAATTTGCCGGGCTGCCTATGGAGTCCACACCA-3'; 5'-AAG GTCTCGCGGGCAAAGGCAGAGTTATTCTTCCCATCCTCAATC AGTG-3'. The TRPV1exCellHalo plasmid is available through Addgene (plasmid #220307). All other TRPV1 constructs were made using either the Quickchange (Agilent) or in vitro assembly (IVA) methods of site-directed mutagenesis as previously described (Mott et al., 2023). For all mutant constructs, our wild-type TRPV1.pcDNA3 and TRPV1-cGFP.pcDNA3 vector was used as a template with either entirely (Quickchange) or partially (IVA) overlapping primers that contain the desired mutation. The full sequence of each completed construct was confirmed using Sanger sequencing.

Ca2+ imaging

Cells were grown on 25-mm coverslips and then incubated for 30 min at room temperature with Fluo-4 (AM; Thermo Fisher Scientific) at a concentration of 3 μ M. The cells were then rinsed with Hepes buffered Ringer's (HBR) solution (in mM: 140 NaCl, 4 KCl, 1.5 MgCl₂, 5 *D*-glucose, 10 HEPES, and 1.8 CaCl₂, and pH adjusted to 7.4 with NaOH) and allowed to rest for another 30 min in HBR at room temperature. The cells were imaged on a

Nikon Eclipse Ti microscope using a 10× objective. For each slip, a brightfield image, a Fluo-4 image, and a 3-min Fluo-4 fluorescence movie with exposures of 100-ms and 0.5-s intervals were captured. During the time sequence, HBR is initially being perfused throughout the chamber. Perfusion is switched to 500 nM capsaicin in HBR at 30 s, and at the 2-min mark, 3 µM ionomycin is added to the chamber. The HBR and 500 nM capsaicin in HBR were perfused into the chamber using open, gravitydriven reservoirs, and 500 µl of 3 µM ionomycin was pipetted into the chamber via micropipette. The data obtained during these experiments were analyzed using ImageJ and Matlab (Mathworks). For each time sequence, ROIs sized 10×10 pixels were placed over approximately seven responding cells where their level of fluorescence was tracked throughout the experiment. For each ROI, the proportion of the maximal fluorescence (F/F_{iono}) that was reached with capsaicin (based on maximal response to ionomycin application) was calculated by removing a baseline offset. The Mann-Whitney U-test was used to determine significance and is given for each indicated comparison.

DRG isolation and transfection with TRPV1exCellHalo

DRG isolation from wild-type C57 mice was performed as described in Stein et al. (2006). Additional supplementation of media with 100 ng/ml nerve growth factor was done to maintain neuronal character as cells were kept for several days. The neuronal culture was transfected with TRPV1exCellHalo-pcDNA3.1 using JetPEI according to the manufacturer's instructions (Polyplus). After 48 h, cells were incubated with 7 μ M Alexa660 HaloTag ligand in complete media for 15 min, which is twice the manufacturer's recommended concentration (Promega). The sample was rinsed once with media and then incubated in complete media for an additional 20 min at 37°C. Coverslips were imaged by epifluorescence with a 40× magnification objective in HBR.



Live cell TRPV1 imaging

Cells expressing either TRPV1-tag constructs or TRPV1exCell-Halo were labeled with their respective click-chemistry fluorescent reagents in an imaging chamber during microscopy imaging experiments. TRPV1-tag pcDNA3.1 constructs were cotransfected into HEK293T/17 cells alongside pAcBac1-NES-R284 at a ratio of 1:2 in the presence of 30 μM Tet3-Butyl for 24-48 h. Subsequent fluorescent labeling with 500 nM sTCO-Cy5 was done in the imaging chamber after exchanging cell culture media for HBR. Cells were incubated in HBR/sTCO-Cy5 labeling solution for 5 min followed by 3 min of continuous rinsing with HBR perfusion. Brightness correlation in Fig. 3 D and Fig. S5 was done with 100× objective, and GFP-filter images were captured via epifluorescence and near-TIRF field excitation of Cy5 dye with Cy5 filter set. Near TIRF means that the laser angle is set to illuminate the sample in an Epi/widefield configuration with reduced background signal because the light passes out to the side of the sample instead of transmitting directly out the top. The raw images shown in Fig. S5 were assessed for mean GFP fluorescence in the indicated ROIs (1-5, mean value shown with each) and compared with maximal Cy5 fluorescence pixel in rectangular ROI for Cy5 channel (1-5, max pixel intensity reported with each). Rectangular ROIs were positioned such that smooth membrane outline fluorescence was captured and bright clusters were avoided. Sampling additional cells across different image fields was complicated by background offsets and strong dependence of intracellular GFP fluorescence signal on the z-axis focal position, which would vary between image fields. Statistics were reported from a linear fit model (fitlm) function in Matlab. R-squared is the coefficient of determination and the P value is returned from an F-test on the linear model with the reported data.

TRPV1exCellHalo was expressed from pcDNA3.1 in HEK293T/17 cells, and cells were replated after 24 h onto poly-D-lysine coated coverslips for imaging. After a minimum of 2 h to allow for cell adhesion, labeling of TRPV1exCellHalo was completed in the imaging chamber by incubating cells with 3 μ M Alexa660 Halo Ligand for 5 min (Promega) in HBR followed by 3 min of continuous rinsing with HBR perfusion.

Single-molecule imaging of TRPV1exCellHalo

Expression of TRPV1exCellHalo in HEK293T/17 cells followed as previously described. After 24 h, cells were transferred from 100-mm dishes to coverslips treated with poly-D-lysine (p7886; Millipore-Sigma) to facilitate adhesion. 4–6 h after replating the cells, we transferred the coverslips to our imaging chamber, labeled TRPV1exCellHalo, and imaged single channels with the following protocol:

- (1) Incubate cells in a chamber with 2 μM Alexa488 HaloTag ligand in HBR for 5 min.
- (2) Rinse cells with continuous perfusion of HBR for 5 min.
- (3) With 100× objective and TIRF imaging locate cells with crowded, yet still resolvable single channels in a cell footprint (smooth fluorescence in cell footprint indicates too high of expression).
- (4) Photobleach cell footprint with the highest laser setting (our system is powered by a 50 mW CW laser and we ordinarily

- maintain a neutral density filter in the laser path; this filter is removed and the laser operated at maximum power) for \sim 5 s. If the EM gain is lowered on the camera, the photobleaching process can be monitored to observe when adequate bleaching is reached.
- (5) Allow recovery of fluorescent TRPV1exCellHalo in the cell footprint for 5 min.
- (6) Acquire one 2.5-s movie (20 Hz) with optimal gain (EM:300 for Andor iXon Life camera) for single-channel imaging and laser intensity modulated to reduce bleaching (we place neutral density filter back into the path and power laser at 50%). This movie is to check if channels diffused sufficiently back into the footprint.
- (7) Begin acquisition of pre-capsaicin movie for 2.5 s (20 Hz).
- (8) Rapidly deliver 5 μM capsaicin solution in HBR into the chamber.
- (9) Wait 15 s after the end of the pre-capsaicin movie to start recording the post-capsaicin movie with identical imaging parameters.

Mock-treated samples follow an identical protocol with HBR added rapidly into the chamber instead of capsaicin solution. Because we wished to see what effect that plating conditions would have on the track displacements of TRPV1exCellHalo, we repeated the mock treatment protocol on cells that were plated on coverslips 1 day before the experiments. Displacement distributions from these mock-treated cells generally exhibited higher mobility than their mock-treated analogues plated on the same day as experiments (see Fig. S6).

Single-molecule TRPV1 tracking

Data collected on moving TRPVlexCellHalo were processed using FIJI (https://imagej.net/software/fiji) by tracking the movement of ion channels within selected ROIs using TrackMate (v3.7.0) (Schindelin et al., 2012; Tinevez et al., 2017). Tracking analysis was performed on movies displaying channel movement just before the application of the capsaicin solution as well as movies displaying channel movement after the capsaicin solution was applied. To test the effects of the capsaicin, two movies were used on the same cell using a prior condition called precapsaicin, and then a capsaicin solution was added in the second.

- (1) Movies obtained from imaging were dragged and dropped into FIJI. Using the freehand selection tool, we selected an ROI over the first frame, taking care to avoid extremely bright clusters of spots in the movie while including as many individual spots as possible.
- (2) After applying the ROI to the movie, we used TrackMate, within FIJI, to track and analyze the movement of the channels throughout the movie. The TrackMate software was accessed from the Tracking option within the drop-down menu under the "Plugins" tab located on the horizontal taskbar across the top of the FIJI window.
- (3) The following settings and configurations were used when performing the tracking analysis using TrackMate:
 - (1) Calibration settings were configured as follows: we set the pixel width and height to 0.160 μm . The voxel depth and time interval were set to 1 μm and 0.052 s,



- respectively. Crop settings were set specific to the ROI of each movie by selecting the movie window and then clicking the "Refresh source" button on this page of TrackMate.
- (2) The LoG detector was selected to apply a Laplacian of Gaussian filter to the image.
- (3) The LoG detector settings were as follows: the estimated blob diameter was set to 0.5 μm and the threshold to 0.5. Additionally, subpixel localization was selected. Clicking the Preview option on this page of TrackMate should display the selection of many fluorescent spots in the movie.
- (4) This page described in detail the abilities and the statistics it will be able to perform.
- (5) The "Initial Thresholding" was set to restrict the sample of spots to be analyzed and tracked. We selected the "Auto" option on this page to consider the spots within the created Mask and chosen ROI.
- (6) The "HyperStack Displayer" View was selected.
- (7) No filters were applied to the spots. Additionally, "Uniform color" was selected as the color setting on this page.
- (8) The "Simple LAP tracker" was selected.
- (9) The Simple LAP Tracker was configured as follows: We specified the maximum linking and gap-closing distance to be 0.5 um. Additionally, the maximum gapclosing frame gap was set to four frames.
- (10) This page summarized the settings we selected to perform tracking.
- (11) No filters were applied to the tracks. The color of the tracks was set based on the "Track displacement."
- (12) The final page summarized the settings we chose for display options. Finally, we selected the "Analysis" button on this page to obtain the complete "Tracking Statistics" data set of the movie we examined.
- (4) All results from the "Tracking Statistics" sheet obtained from FIJI and TrackMate were selected and copied into an Excel spreadsheet for further analysis.
 - (1) In Excel, the data were sorted in descending order based on column B, "Number Spots." We selected only the data with the value of Number Spots >5 spots to only examine tracks of 5 spots or longer. The data of 5 number spots and higher was pasted in another sheet within the file.
- (5) We chose to analyze the specific tracking statistic, Track Displacement, using MATLAB (https://www.Mathworks.com). Using MATLAB, we created figures representing trials with capsaicin or mock treatment in cells that were plated the same day or a day earlier.
 - (1) For each experiment, we used MATLAB to create a figure displaying cumulative density histograms for the pre- and post-capsaicin solution conditions. The Track Displacement data was imported into MATLAB and normalized from 0 to one to make the comparison between the different pre- and post-capsaicin solution histograms possible. Both histograms were overlayed within one figure to visualize overall shifts in data.
 - (2) We extracted the upper quartile values of both pre- and post-capsaicin solution data sets. We used MATLAB to

- sort and index the data sets, before multiplying the data set by 0.75 and taking the floor value as the index number to obtain the upper quartile displacement value.
- (3) With two upper quartile values per experiment (before and after capsaicin). We plotted the pairwise comparisons in each experimental set.
- (4) Finally, in order to statistically analyze the potential effects of capsaicin activation, we performed a MATLAB two-sample Student's t test between the pre- and post-capsaicin solution data sets for all experimental conditions.

Western blotting

For Western blotting, cells were incubated for 48 h at 37°C following transfection. The media was replaced after 24 h. Cells were collected from culture plates and washed with PBS before lysing with the following steps. The protein fraction was isolated by lysing the cells using Lysis buffer + HALT (25 mM Tris-HCL, 150 mM NaCl, 1 mM EDTA, 1% Triton, and HALT [Thermo Fisher Scientific]) and in accordance with the Western blot protocol used by Zagotta et al. (2016). The lysate was loaded into a PAGE SDS precast gel and was run at 200 V for 40 min. Using a BOLT wet transfer setup at 20 V for 60 min, the protein was transferred from the PAGE gel to a PVDF membrane (Thermo Fisher Scientific). The membrane was then preblocked in 5% skim milk in PBST for 1 h while shaking. The primary antibody (anti-GFP: TP401, Torrey Pines Biolab) was then added 1:5,000 directly to the blocking solution, and the membrane was covered and placed overnight on the shaker. The following morning, the membrane was rinsed with PBST and fresh 5% skim milk in PBST was added to the membrane. The secondary IR fluorescencelabeled antibody was added 1:15,000 to the blocking solution. Following a 45-min incubation in this solution while shaking, the membrane was imaged on a LI-COR imager with 700 and 800 nm wavelengths (LI-COR Biosciences).

PAGE gel-shift assay

Cell lysate from HEK293T/17 cells expressing the TRPV1-tag construct of interest with Tet3-Butyl as the ncAA was prepared as described in the previous section. Expression of sfGFP-N150tag with Tet3-Butyl was done by cotransfecting HEK293T/ 17 cells with pAcBac1-sfGFP-N150tag and pAcBac1-NES-R284 at an 8:1 ratio and supplementing the cell culture media with 30 μM Tet3-Butyl. Cell lysis buffer with 1% TritonX100 was used as described for Western blotting in Fig. S4 A. To each "+ sTCO-PEG5K" sample, we added sTCO-PEG-5K for a final concentration of 100 µM. The reaction was quenched after 10 min or variable time by adding Tet3-Butyl to a final concentration of 3 mM. Treated lysate samples were loaded onto a PAGE precast gel and run at 200 V for 40 min. RIPA buffer was used to lyse cells for the gel shift assay in Fig. S4 B. Laemmli buffer was added to each sample for a 1× concentration and then samples were loaded into a 12% SDS-PAGE gel and ran between 150 and 200 V until the dye front ran off the gel. Western blotting to identify GFP-construct bands was completed as described above beginning with transfer to the PVDF membrane and using either the GFP antibody (TP401: see above) or anti-FLAG for Fig. S4 B.



Co-immunoprecipitation

Immunoprecipitation of TRPV1exCellHalo was done with Halo-Trap resin (catalog: ota) and as described in packaging instruction (Proteintech) with the following changes: (1) For $\sim\!\!6\times10^6$ cells, we used 500 μl lysis buffer of identical composition as used in the previously described Western blot method. (2) Wash buffer contains (in mM) 100 Tris pH 7.6, 150 NaCl, 0.1 EDTA, 1 mg/ml BSA, and 1× HALT. (3) Final wash step was done in wash buffer without BSA and HALT. (4) Elution for gel loading is in 40 μL wash buffer and 1× SDS loading buffer (250 mM Tris pH 6.8, 8% SDS, 50% glycerol, 0.1% bromophenol blue), and 10% β -mercaptoethanol. (5) Sample is not boiled prior to PAGE gel loading. Western blot analysis of TRPV1-cGFP pulldown fraction was done as previously described.

Online supplemental material

Fig. S1 shows protein expression of TRPV1-tag constructs. Fig. S2 shows the expression of TRPV1-V469tag-GFP in the absence of pyl-RNA synthetase/tRNA^{pyl} (pNEU). Fig. S3 shows TRPV1-T468tag-GFP and TRPV1-K464tag-GFP negative control. Fig. S4 shows Western blots of gel-shift assays with sfGFP-N150tag/Tet3-Butyl and sTCO-PEG5K. Fig. S5 shows brightness correlation analysis of TRPV1-T468tag/Tet3-Bu-GFP expressing cells and labeled with sTCO-Cy5. Fig. S6 shows cumulative density histograms of track displacements before and after capsaicin treatment. Video 1 shows image sequence capture under precapsaicin conditions in our capsaicin treatment sample.

Results

Missense variants occur more frequently in intracellular and extracellular loops

We performed a bioinformatic analysis on human genomic data to replace an amino acid residue with an ncAA on the extracellular side of TRPV1, where it is unlikely to impact channel function. As the number of human genome sequences added to genomic variant databases increases, a growing body of research is applying this new-found sequence knowledge to protein structure and function (Plante et al., 2019; Bai et al., 2021). Previous studies of genomic missense variants occurring in the human population reveal a correlation pattern between buried side chains in the TRPV1 N-terminus and a reduction in the number of variants for the corresponding residue (Mott et al., 2023). We have extended this pattern analysis into the transmembrane region, which contains conserved sequences assigned to the transmembrane segments, S1-S6, in TRPV1 and connecting segments of reduced conservation that correspond to internal and external loops (Fig. 2 A, top). We narrowed down potential sites for ncAA insertion in the external loops of TRPV1 by implementing an algorithm to count the absolute number of missense variants at each residue and plotted these data against the secondary structure of the transmembrane region (Fig. 2 A). The most salient feature of these plotted data is the increased occurrence of missense variants in regions connecting the transmembrane segments, precisely where we intend to replace an amino acid with a ncAA. However, the missense variant counts also show great variability within the extracellular and

intracellular loops. Informed by the results of our bioinformatics study, we empirically determined the optimal placement of a ncAA, LysZ, or TCO, at variant sites along the loop connecting S1 and S2, where the perturbation is expected to minimally affect the pore gating mechanism (Fig. 2 B, yellow). Success of the ncAA insertion in tag amber codon sites of TRPV1 was evaluated by Ca2+ imaging experiments in TRPV1-tag-GFP expressing HEK293T/17 cells. Previous work with TRPV1-Q169tag-YFP allowed us to evaluate TCO insertion with the pNEU amber codon suppression at novel sites in the S1-S2 loop (Zagotta et al., 2016). In comparison to background activity with cells that showed no appreciable change in fluorescence on exposure to 500 nM capsaicin, TRPV1-Q169tag expression in cells incubated with the TCO incurred a relatively significant response (Fig. 2 C, Q169tag black versus purple markers). The difference in capsaicin response between transfected cells and background is shared by TRPV1 constructs with tag codons replacing the following amino acids in the S1-S2 loop: K464, T468, and V469 (Fig. 2 C). A notable result of our Ca2+ imaging experiments is the robust TRPV1-V469tag response obtained in the absence of a ncAA. Without co-expression of pNEU and supplementation with TCO there is no appreciable expression of full-length TRPV1-K464tag or TRPV1-T468tag, and this is in stark contrast to the expression product of TRPV1-V469tag visible in a Western blot (Fig. S1 A). We investigated the properties of TRPV1-V469tag further by varying the circumstances under which the channel is expressed and determined that the TRPV1-V469tag-GFP construct requires neither co-expression of the pNEU plasmid nor supplementation with a ncAA to achieve appreciable expression (Fig. S1 B and Fig. S2). Moreover, even in the presence of the suppression plasmid, pNEU, TRPV1-K464tag-GFP, and TRPV1-T468tag still require the ncAA (TCO) to form functional channels in our Ca2+ imaging experiments (Fig. S3). We summarize our results for scanning potential ncAA insertion sites in the S1-S2 loop in Fig. 2 D, which includes a sequence conservation alignment and illustrates the stepwise procedure we used from identifying missense variants in the human gnomAD sequence ("Variant") to experimentally assess TRPV1-tag-ncAA-GFP functional channels.

Cell surface fluorescence labeling of TRPV1 at single residues

The reduced expression level of previously characterized TRPV1-Q169tag-GFP construct with the pNEU support plasmid (Western blot band in TRPV1-GFP lane compared with the band in Q169tag-GFP of Fig. S1) motivated us to test alternative amber codon suppression tools. We found that the tetrazine-based ncAA insertion tools provided through the GCE4All Research Center were the optimal method for ncAA insertion and chemical modification of K464 and T468 (Jang et al., 2020; Ryan et al., 2022). In HEK293T/17 cells that express TRPV1 tag mutations at K464 and T468, partial reactivity of the Tet3-Butyl ncAA with an sTCO-PEG5K substrate could be observed through an up-shift in the molecular weight bands of TRPV1-tag-GFP (Fig. 3 A, lanes 1 and 5). Repeating the Ca²⁺ imaging study of TRPV1-T468tag-GFP with required GCE4All support plasmid (pAcBac1) and supplementation with ncAA Tet3-Butyl demonstrated a robust activation of channels in response to capsaicin (Fig. 3 B and Fig. S4 C). An important aspect of implementing amber codon



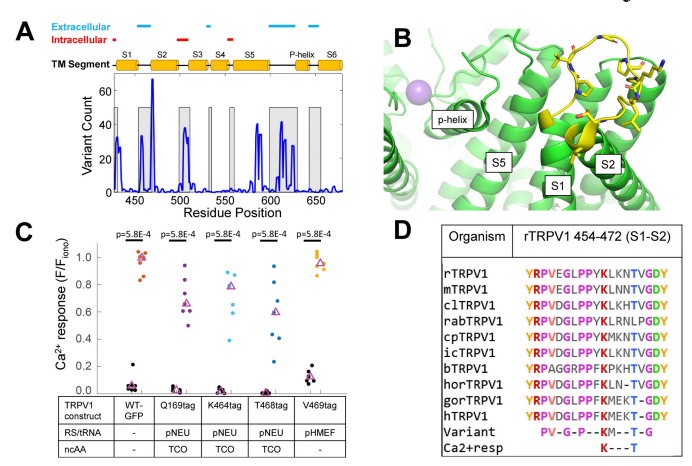


Figure 2. **Insertion of ncAAs between transmembrane segments 1 and 2 of TRPV1. (A)** Missense variants encoded in the TM domains of human TRPV1 (residues 427–680) as annotated in the gnomAD ver. 2.1.1. The blue line is the absolute variant count for the given residue filtered across three residues with a boxcar kernel. TM segments 1–6 are illustrated above the plot (yellow cylinders) and connecting loops are shown as grey regions within the plot. Extracellular (blue) and intracellular (red) loop regions above TM segments are potential sites for ncAA insertion. (B) A reconstructed structural model of one TRPV1 subunit from cryo-EM data (PDB accession no. 7L2H; green) reveals details of the extracellular loop between transmembrane segments 1 and 2 (yellow). (C) Ca²⁺ imaging experiments to determine the successful expression of full-length TRPV1 constructs containing tag codons in the S1–S2 extracellular loop. RNA-synthetase/tRNA plasmid and type of ncAA used are given at the foot of the plot. The median value of the set is indicated with a purple triangle. Significant differences between Ca²⁺ imaging responses from non-responding cells (black) and cells with responses to 500 nM capsaicin (colored) within the same imaging experiment were determined by Mann–Whitney U-test. (D) Alignment of rat TRPV1 (rTRPV1) sequence (454–472; S1–S2 extracellular loop) against homologous sequences from several mammals (m: mouse [UniProt Q704Y3]; cl: dog [UniProt Q697L1]; rab: rabbit [UniProt Q6RX08]; cp: guinea pig [UniProt Q6RS63]; ic: 13-lined ground squirrel [UniProt I3LZN5]; b: bovine [UniProt E1BGM0]; hor: equine [UniProt A0A3Q2H4H5]; gor: gorilla [UniProt G3R334]; h: human [UniProt Q8NER1]). Variant positions identified in the gnomAD 2.1.1. Database are listed below the conservation alignments with P456, V457, G459, P461, K464, T468, and G470 standing out as highly conserved residues (conservation >90%). Our first round of calcium imaging with P456tag, V457tag, K464tag, T468tag, and V469tag returned functional, ncAA-depend

suppression tools for surface labeling in pulse quench experiments is that the chemical reaction rate between the ncAA and the chemical label must be rapid. We confirmed the rapid rate of labeling with the GCE4All system by expressing super-folder (sf) GFP as a tag construct at position N150 in HEK293T/17 cells before lysing these and subjecting the free Tet3-sfGFP¹⁵⁰ to pulsed labeling with sTCO-PEG5K for varying intervals of time. Our Western blots of pulsed sTCO-PEG5K showed that maximal labeling of the Tet3-sfGFP¹⁵⁰ is achieved in 3 min (Fig. S4 A, compare lane 2 [3 min] with lane 3 [10 min]). Our method for lysing HEK293T/17 cells and extracting TRPV1 proteins is, at least, partly the reason why there is incomplete labeling of Tet3-sfGFP¹⁵⁰ with sTCO-PEG5K. Under optimal sfGFP extraction conditions using RIPA buffer (see Materials and methods), complete labeling of Tet3-sfGFP¹⁵⁰ is possible with sTCO-PEG5K

(Fig. S4 B). Ultimately, for the extracellular labeling of TRPV1-tag ncAA constructs to have any practical use, reactivity with the sTCO-Cy5 dye needs to be sufficient for fluorescence imaging. To examine the threshold of reaction concentration and time needed for acceptable labeling of TRPV1-K464tag/T468tag-GFP channels, we expressed the construct in HEK293T/17 cells for 48 h while in the presence of 30 μ M Tet3-Butyl. After replating the cells on coverslips for imaging and mounting the coverslips in a chamber, we applied a labeling solution of 500 nM sTCO-Cy5 for 5 min before 3 min of washing. Cells expressing TRPV1-T468tag-GFP as determined by GFP fluorescence were confirmed to have surface labeling by sTCO-Cy5 (Fig. 3 C). We were able to establish that the amount of sTCO-Cy5 label increased with expression of TRPV1-T468tag/Tet3-Bu-GFP by correlating the GFP fluorescence signal with Cy5 fluorescence on

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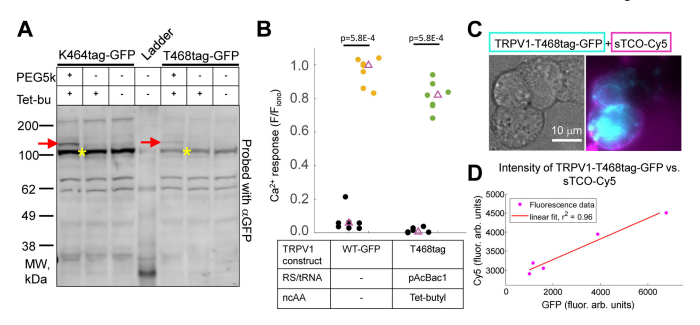


Figure 3. **TRPV1-tag construct expression with GCE4All-Tetv3.0 system. (A)** Western blot of TRPV1-K464tag-GFP and TRPV1-T468tag-GFP with pAcBac1 and Tet3-Butyl ncAA. Addition of sTCO-PEG5k (PEG5k) used to test access to tetrazine ncAA in TRPV1-tag target. Excess Tet3-Butyl (Tet-bu) is added to quench sTCO-PEG5k at the 10 min point to prevent off-target labeling with PEG5K. TRPV1-tag-GFP molecular weight is estimated as ~120 kD (see yellow asterisk) and TRPV1-tag-GFP/sTCO-PEG5K would be upshifted to ~125 kD (see red arrow). **(B)** Ca²⁺ imaging experiment to test the functionality of TRPV1-T468tag-GFP expressed with GCE4All-Tetv3.0. Median value of the set is indicated with a purple triangle. Significant difference was observed between the Ca²⁺ response evoked by 500 nM capsaicin in TRPV1-T468tag-GFP cells (green) and non-responsive cells (black) in the same imaging experiment (Wilcoxon rank sum; P < 0.001). Only results of Ca²⁺ imaging with the T468tag construct are shown because labeling experiments with this construct and sTCO-Cy5 are demonstrated in panels C and D. **(C)** HEK293T/17 cells expressing TRPV1-T468tag/Tet3-Bu-GFP (cyan) are labeled with sTCO-Cy5 (magenta). **(D)** Correlation of GFP brightness in the image of TRPV1-T468tag/Tet3-Bu-GFP expressing cells that are labeled with 500 nM sTCO-Cy5 for 5 min as described in methods. Five cells expressing various levels of the TRPV1-T468tag/Tet3-Bu-GFP construct as determined by GFP intensity show correlated labeling with sTCO-Cy5 (linear fit, r² = 0.96; P value = 0.0034; see Fig. S5 and Materials and methods for description of statistics). Source data are available for this figure: SourceData F3.

the cell surface (Fig. 3 D and Fig. S5). TRPV1-K464tag/Tet3-Bu-GFP showed inconsistent sTCO-Cy5 labeling patterns (data not shown), and we elected to focus our continued efforts on the TRPV1-T468tag-GFP construct. Additional verification of the expression, function, and labeling properties of TRPV1-T468tag/Tet3-Bu-GFP are reported by Koh and coworkers (Koh et al., 2024, *Preprint*).

Cell surface labeling of TRPV1 with circularly permutated HaloTag

Our amber codon suppression tools for labeling TRPV1 permit a small footprint labeling scheme with minimal perturbation to channel function. However, the amber codon suppression method for protein expression has much lower efficiency than a construct with the 20 naturally occurring amino acids (compare lanes 1 and 2 of Fig. S1 A). To complement our single residue surface labeling approach with a high expression TRPV1 product, we engineered an insertion point for the circularly permutated (cp) version of HaloTag (Deo et al., 2021). Our TRPV1exCellHalo construct contains a modified turret domain between transmembrane domain S5 and the pore helix with adequate space for the cpHaloTag to be accessible from the extracellular space in the fully assembled TRPV1 tetramer (Fig. 4 A). Ca²⁺ imaging experiments of TRPV1-exCellHalo showed robust responses to 500 nM capsaicin (Fig. 4 B). Evaluation of TRPV1exCellHalo capsaicin sensitivity by dose-response electrophysiology recordings revealed only a mild increased gain of function (Fig. 4 C). Apart from the non-significant increase in capsaicin

sensitivity, TRPV1-exCellHalo appears to behave like the wild-type counterpart. To assess the labeling properties of the cpHaloTag domain on the extracellular side of TRPV1, we incubated HEK293T/17 cells that expressed the TRPV1-exCellHalo construct with 3.5 µM Alexa660-HaloTag for 5 min. Surface labeling with these conditions was sufficient for clear visualization of the plasma membrane on cells that also responded to capsaicin in a Ca²⁺ imaging experiment (Fig. 4 D). Since the TRPV1exCellHalo construct was designed with the purpose of using it in cell culture and primary cell lines, we wished to test if the subunits containing cpHaloTag were able to co-assemble with intact TRPV1 containing a C-terminal GFP (TRPV1-cGFP). In an immunoprecipitation study of TRPV1-exCellHalo co-expressed with TRPV1-cGFP, we used a HaloTag antibody pulldown protocol and probed for TRPV1-cGFP with a GFP antibody. By pulling down the TRPV1-exCellHalo subunits of our co-expression lysate, we obtained an enrichment of TRPV1-cGFP in our Western blot, which was not the case for lysate derived from cells expressing only TRPV1-cGFP (Fig. 4 E). Based on these results, we reason that exogenously expressed TRPV1-exCellHalo subunits may freely co-assemble with endogenously expressed TRPV1 in primary neurons.

Cell surface tracking of TRPV1exCellHalo

The prospect of using a rapid labeling, photostable, and exclusively surface-labeled TRPV1 construct appealed to our group's interest in tracking cell surface-bound receptors. Our previous study on activity-dependent mobility of cell surface TRPV1 relied



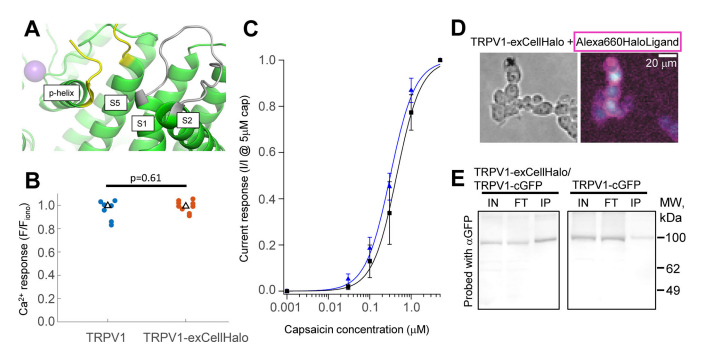


Figure 4. **Expression and functional profile of TRPV1-exCellHalo. (A)** Reconstructed model of the extracellular region on a TRPV1 subunit (PDB accession no. 7L2H). The loop between TM S1 and S2 used for ncAA insertion is shown in grey. Yellow termini demarcate the disordered region between TM segment 5 and the pore helix where circularly permutated HaloTag is inserted. **(B)** Ca²⁺ imaging experiment in HEK293T/17 cells comparing responses of wild-type TRPV1 against TRPV1-exCellHalo when activated by 500 nM capsaicin. Statistical comparison determined by Mann–Whitney U-test. **(C)** Dose response experiment of wild-type TRPV1 (n = 4; $K_{1/2} = 0.53 \pm 0.28$) and TRPV1-exCellHalo (n = 7; $K_{1/2} = 0.34 \pm 0.19$). These $K_{1/2}$ values are not significantly different as determined by Student's two-tailed t test (P = 0.66). **(D)** Left: DIC image ($10 \times magnification$) of HEK293T/17 cells transfected with TRPV1-exCellHalo. Right: Fluorescence Ca²⁺-imaging experiment of TRPV1-exCellHalo expressing cells shown in the left panel with Fluo4 following activation with 500 nM capsaicin (cyan). After Ca²⁺ imaging experiment, surface labeling is accomplished with the incubation of HaloTag ligand Alexa660 (magenta) for 3 min. **(E)** Western blot of immune precipitation with anti-HaloTag antibody (Chemtek). Cell lysate is incubated with anti-HaloTag resin and the blot is probed with anti-GFP antibody. Two conditions tested are co-expression of TRPV1-exCellHalo/TRPV1-cGFP or TRPV1-cGFP alone (IN: input; FT: flow through; IP: immunoprecipitated). Source data are available for this figure: SourceData F4.

on a genetically encoded GFP fused to an intracellular domain of the channel (Senning and Gordon, 2015). TRPV1exCellHalo expressed on the cell surface is exclusively labeled by cell impermeable dyes while the intracellular fraction is not modified, and the rapid labeling process ensures that labeled TRPV1exCellHalo remains on the surface during the experiment. We devised a similar experiment to our previous single molecule study of TRPV1-GFP described in Fig. S6 of Senning and Gordon (2015), evaluating the changes to channel mobility after activation by capsaicin. Because our previous TIRF experiments with TRPV1-GFP could not guarantee a uniform channel population on the cell surface, we had to resort to an elaborate analysis of the channel dynamics. Here, we were able to measure the displacement of individual TRPV1exCellHalo on the surface of HEK293T/17 cells before and after cells are treated with capsaicin and carry out a direct comparison between the distribution of displacements under the different conditions (Fig. 5, A and B). Total displacements of TRPV1exCellHalo tracks recorded in a 2.5s movie (Video 1) prior to the rapid addition of 5 µM capsaicin have a distinctly faster cumulative density histogram profile (blue) compared with the histogram of tracks after capsaicin treatment (red) and recorded 15 s later (Fig. 5 C). For each data set, we compared the upper-quartile displacement value (0.75; dashed black line in Fig. 5 C) to demonstrate the changes in displacements occurring with capsaicin or mock treatments

(Fig. 5 D and Fig. S6). A notable aspect of the mock experiments done on the same day of plating was that pre- and post-treatment upper-quartile displacements were lower than in the capsaicin set of experiments, and we were able to see these values increase to those observed in capsaicin experiments by plating the cells on coverslips 1 day before the experiment. Regardless of what value ranges the displacements showed initially in our mock experiments, we did not observe the significant decrease in displacements observed in our capsaicin-treated experiments. Our study of single molecule TRPV1exCellHalo mobility could, therefore, substantiate our earlier study that activation of TRPV1 channels by capsaicin decreases the lateral movement of the channels on the cell surface. To assess the surface expression properties of TRPV1exCellHalo in a more physiological preparation, future studies will rely on the expression of the channel in isolated DRG neurons, which we have shown is possible along with satisfactory labeling by HaloTag ligands (Fig. 5 E).

Discussion

There is a critical need to improve extracellular labeling methods for studies of channel trafficking and cell surface expression dynamics. Extensive use of labeling schemes ranging from immunofluorescent methods to FlaSH/ReASH demonstrates the technological breadth of techniques used in these studies.



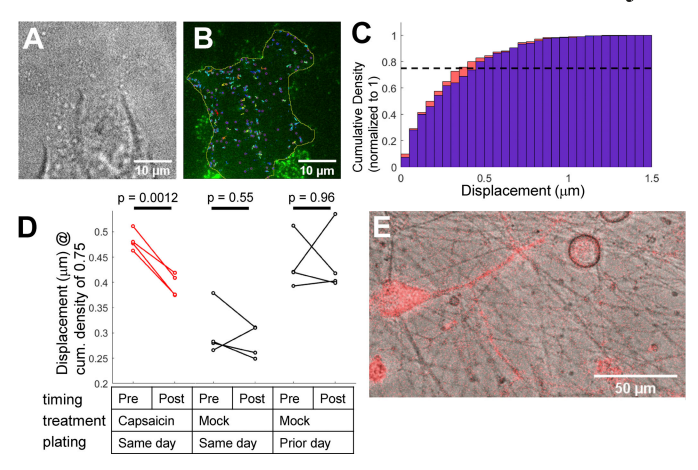


Figure 5. **Reduced mobility of single TRPV1exCellHalo channels after capsaicin treatment. (A)** DIC image of HEK293T/17 cells that transiently express TRPV1exCellHalo. **(B)** TIRF microscopy image of the cell is shown in panel A after labeling TRPV1exCellHalo with Alexa488 HaloTag ligand. The image shown is the first frame of a 2.5-s movie acquired after pre-capsaicin treatment that reveals individual channels diffusing laterally across the cell surface (See Video 1). Overlaid onto the image is the ROI (yellow) to define the area in which ion channels are tracked and individual tracks obtained during the movie. **(C)** Cumulative density histogram of individual channel track displacements. The total number of tracks is normalized to 1.0 in order to compare displacement distributions from tracks in a movie acquired pre-capsaicin (blue; tracks of panel B and Video 1) to tracks acquired in a second (red) movie 15 s after rapid buffer exchange with a 5 μM capsaicin solution. Dashed, black line guides the eye to the upper-quartile (0.75) value in each distribution. **(D)** Quantitative comparison between track displacements from pre-capsaicin movies and post-capsaicin movies are made by evaluating the difference in the upper-quartile (0.75) displacement value (see the full set of distributions in Fig. S6). Mock treatment experiments with rapid buffer exchange that contains no capsaicin and day in which the cells are replated onto coverslips for the experiment are noted in table. Statistical significance between pre- and post-capsaicin treatment is determined by paired Student's t test. **(E)** Transient expression of TRPV1exCellHalo in isolated mouse DRG culture followed by labeling with Alexa660 HaloTag Ligand.

However, experiments that examine protein dynamics are fundamentally constrained by the limitations imposed by the available labeling technology. For instance, in the case of immunofluorescent methods that target genetically encoded epitopes, concern arises due to the perturbation imposed by the large footprint of the antibody. On the other hand, the FlaSH and Re-ASH system, although sound in principle, is subject to off-target labeling, is sold as a cell-permeant reagent, and has longer incubation times (~30-60 min according to a manufacturer's kit available through Thermo Fisher Scientific) (Stroffekova et al., 2001; Specht et al., 2017). Many alternative strategies exist, and yet the benchmarks for technique optimization are unchanged: use a minimally perturbing labeling scheme that is specific and rapid. We therefore sought to address these needs by setting two goals: (1) identify sites in extracellular regions of the ion channel TRPV1 that would tolerate perturbations from sequence substitutions and insertions without critically impacting function, and (2) be able to implement pulse-chase time course

experiments through a rapid labeling scheme. We also relied on a two-pronged strategy for labeling techniques to avert perceived or apparent weaknesses in one approach by exploring a complementary technique in parallel.

Our first approach implemented the amber codon suppression technique of replacing a single amino acid in the extracellular loop between S1 and S2, and we selected optimal ncAA insertion points by exploiting recently collected human genomic data available through gnomAD. We surveyed the extracellular loops, which connect transmembrane segments of the channel, and were able to refine our selection of tag sites to experimentally test the feasibility of inserting the ncAA at a handful of positions between S1 and S2. Ca²⁺ imaging experiments used to test these sites revealed that K464, T468, and V469 could be substituted with the amber codon for protein termination, tag. However, results obtained under various expression conditions indicated that V469tag did not require the supplementary RNA synthetase and tRNA components for efficient translation of a

Fluorescence labeling of cell surface TRPV1



full-length and functional TRPV1 channel. Although our experiments allowed us to eliminate potential ncAA insertion sites that were identified in our missense variant screen, we did not expect that V469tag would show such prominent readthrough, even when compared to supplementation with the pNEU plasmid and different ncAAs (Fig. S1 B). Translational readthrough at the tag amber codon (+1, +2, and +3 positions) occurs as a sporadic event in eukaryotic cells and is partially determined by the flanking sequence nucleotides and interactions with the near-cognate tRNA or translational machinery (Dabrowski et al., 2015). Downstream nucleotide preference for a "c" at the +4 position is a prominent marker for readthrough at stop codons "taa" and "tga," but for tag, a notable exception is made where multiple "g"s may appear at positions +4,+5, and +6 in gag-pol coding regions of mammalian type C retroviruses (Feng et al., 1992; Alam et al., 1999). In our TRPV1-V469tag sequence, the nucleotide corresponding to the two codons after tag is "ggggac." Perhaps more relevant to our task of inserting an ncAA, the Bultmann group devised an online tool (iPASS) to assess tolerance of the flanking of DNA sequences to either side of a potential tag site (Bartoschek et al., 2021). The iPASS tool returns a simple test score wherein a threshold over 1.0 indicates sufficiency for ncAA insertion. Both permissive sites we isolated, K464 and T468, returned scores of 1.16 and 0.87, respectively. However, the TRPV1-V469tag mutation and surrounding sequence gave an iPASS score of 0.05.

In a similar bid as implemented with ncAA incorporation, we attempted to insert the cpHaloTag domain at specific sites in the loop between S1 and S2 that coincided with increased missense variants. However, these constructs failed to express on the surface of HEK293T/17 cells. We could not readily discern the reason for this based on our experimental results, but a literature search for extracellular loop fusion proteins uncovered a recent study in which similar placement of genetically encoded fluorescent proteins in the ABCC6 integral membrane protein was also problematic, possibly due to short extracellular loops that would lead to folding or trafficking deficits if large polypeptide tracts were introduced (Szeri et al., 2021). Based on unsuccessful attempts at finding an insertion site between the S1 and S2 of TRPV1, we sought an alternate location and added the cpHaloTag in the "turret domain" of TRPV1 between S5 and the pore-helix (Cui et al., 2012). Concerns over whether a fusion protein above the pore domain would impact function were laid to rest when our Ca²⁺ imaging assays and e-phys dose-response experiments of the TRPV1exCellHalo construct nearly overlapped with the wild-type function.

Once the ncAA insertion site was successfully determined by testing TRPVI functionality, we addressed the second goal of our project to ask how we would improve the click-chemistry reaction rate between the ncAA and ligand reporter, such as a fluorescent dye molecule. Our experience with the pNEU vector using TCO as the ncAA was complicated by inconsistent yield during the labeling stage with the tetrazine-fluorophore. To address the erratic properties of the pNEU/TCO system, we switched to the GCE4All amber codon suppression system that relies on a tetrazine-based ncAA and sTCO ligand (Jang et al., 2020). Both our in vitro labeling reaction with an sTCO-PEG5K

ligand and on cell labeling with sTCO-Cy5 could be performed within 5 min, which is well within the time needed for pulse chase experiments that would permit measurement of membrane trafficking events (Fig. 3). Switching to the GCE4All system also partially assuaged our concern over whether an excess burden of partial or abortive translation products may interfere in TRPV1 trafficking or membrane localization studies. There is at least some qualitative evidence from our Western blots with pAcBac1 that shows improvement in the ratio of full-length TRPV1 product (~120 kD) relative to partial-length products (~62 kD) when compared with pNEU (see Q169tag in Fig. S1 A and K464tag/T468tag in Fig. 3 A). Moreover, the study by Koh and coworkers made extensive use of the TRPV1-T468tag construct in a trafficking study and characterized its functional properties as well (Koh et al., 2024, Preprint).

The rapid ~5 min reaction rate of HaloTag with its chloroalkane substrate is a valued property of this genetically coded click-chemistry product (Jonker et al., 2020). However, it was not certain whether our fusion of the circularly permuted HaloTag in the TRPV1exCellHalo construct would retain labeling with fluorescent ligands. Accessibility of the substrate binding site had been a concern given the permutated sequence of cpHaloTag and a position in our TRPV1 construct that imposed a crowded configuration above the pore domain. In our hands, we could label surface TRPV1exCellHalo with Alexa660-HaloLigand in 5 min, which fulfilled our criteria for a rapid labeling step. Although our fluorescent HaloLigand assay was successful, we were not able to successfully use the HaloLink resin product (Promega) as a pull-down matrix in a co-immunoprecipitation assay for heteromeric assembly of TRPV1exCellHalo and TRPV1cGFP (data not shown). An alternative co-immunoprecipitation assay with IgG specific to HaloTag was still able to recognize the cpHaloTag and could confirm our hypothesis that TRPV1ex-CellHalo will heteromerically assemble with TRPV1-cGFP and presumably also wild-type TRPV1 subunits, and we speculate that the enzymatic site in TRPV1exCellHalo is occluded from matrix in the HaloLink product.

With the successful insertion of cpHaloTag into TRPV1 and intact capsaicin functionality, we moved forward with an experiment to test whether the mobility of TRPV1exCellHalo is changed by channel activation. Our experiment is a reprise of our study done with TRPV1-GFP in Senning and Gordon (2015). However, we can now track the subset of channels only expressed on the cell surface without signal contamination from intracellular pools. Our results with TRPVexCellHalo and Alexa488 HaloTag ligand reaffirmed the results from our earlier study: capsaicin activation of TRPV1exCellHalo in Ca²⁺-containing buffer causes a reduction in the mobility of the channel. The mechanism for this activity-dependent decrease in mobility remains elusive, and we speculate that possibilities range from increased interactions between TRPV1 and the cytoskeleton that are facilitated by Ca2+ to internalization of the channel through endocytosis. An interesting observation in our current study was that the TRPV1exCellHalo channels exhibited different degrees of mobility based on how much time had passed since the cells were replated from 100-mm dishes to poly-D-lysine coated coverslips. Evidently, the substrate under the HEK293T/17 cells



would appear to influence the overall properties of channel mobility. Although there may be variability observed in channel displacement properties of cells that are replated on the same day (capsaicin and mock-treated trials replated same day), the displacement shifts to indicate greater mobility when cells remain on the same substrate for longer times (mock trial on day after replating). That TRPV1 mobility could be sensitive to the underlying substrate of the cell is worth investigating further, especially considering that we do not know if this mechanism is mediated by direct contact with the extracellular matrix or adhesion factors within the cell.

Taken together, TRPV1-tag/ncAA and TRPV1exCellHalo constructs form a comprehensive set of tools to investigate trafficking properties and cell surface expression of TRPV1. With only a single residue in TRPV1 converted to an ncAA in the amber codon suppression construct TRPV1-tag, we minimize the labeling perturbation to the site of an amino acid sidechain. The tradeoff for a minimal label is the limited expression efficiency of the construct. Our second construct, TRPV1exCellHalo, takes advantage of genetically encoding the HaloTag domain in an extracellular loop at the expense of a larger molecular perturbation, but this construct comes with distinct advantages in expression efficiency while maintaining rapid labeling and near wild-type function. It is also noteworthy that the intracellular domains of TRPV1exCellHalo are unchanged and should be capable of sustaining all intracellular interactions. Moreover, the effect on cellular transport and expression by including the extracellular domain in TRPV1 can be studied and contrasted against the minimally perturbed TRPV1-tag/ncAA channel.

Data availability

The data underlying Fig. 2, A and D, were sourced from the gnomAD project (https://gnomad.broadinstitute.org/gene/ENSG00000196689?dataset=gnomad_r2_1) and Uniport (accession nos. O35433, Q704Y3, Q697L1, Q6RX08, Q6R5A3, I3LZN5, E1BGMO, A0A3Q2H4H5, G3R334, and Q8NER1), respectively. All other data are available from the corresponding author upon reasonable request.

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Writing—review & editing, A.J. Eddins: Data curation, Methodology, Writing—review & editing, R.A. Mehl: Conceptualization, Funding acquisition, Resources, E.N. Senning: Conceptualization, Data curation, Formal analysis, Funding acquisition, Investigation, Methodology, Project administration, Resources, Software, Supervision, Validation, Visualization, Writing—original draft, Writing—review & editing.

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Supplemental material

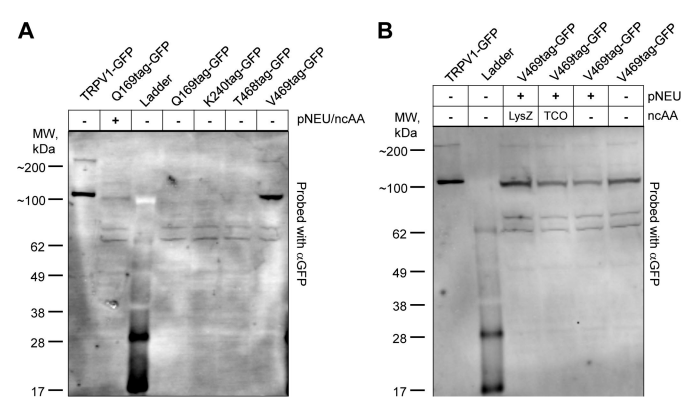


Figure S1. **Protein expression of TRPV1-tag constructs. (A)** Testing of construct TRPV1-Q169tag in the presence and absence of RNA synthetase/tRNA plasmid (pNEU) + ncAA (TCO). Lanes 5, 6, and 7 contain expression products of TRPV1-K240tag, T468tag, and V469tag without pNEU. All lanes are probed with GFP antibody. **(B)** Testing of TRPV1-V469 in the presence and absence of pNEU and two different ncAAs (TCO or LysZ). Lanes are probed with the GFP antibody. Source data are available for this figure: SourceData FS1.



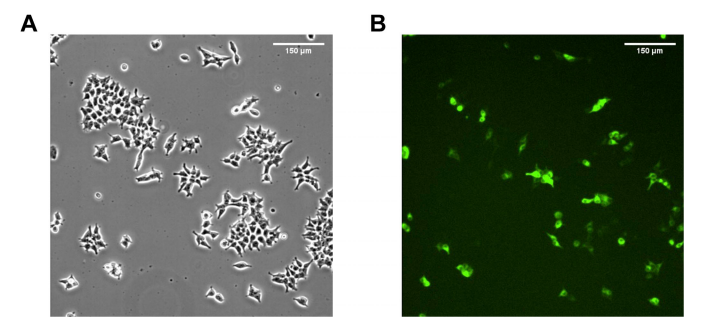


Figure S2. Expression of TRPV1-V469tag-GFP in the absence of pyl-RNA synthetase/tRNA^{pyl} (pNEU). (A) DIC image of HEK293T cells expressing TRPV1-V469tag-GFP. (B) Green fluorescence image of cells shown in panel A.

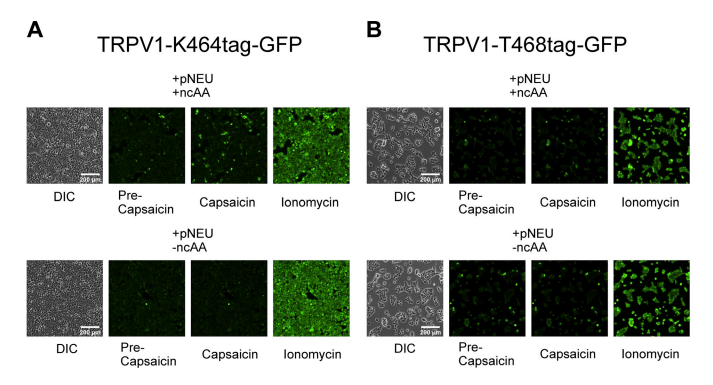


Figure S3. **TRPV1-T468tag-GFP and TRPV1-K464tag-GFP negative control.** Expression of functional TRPV1 channels is qualitatively absent when assessing Ca^{2+} imaging responses to 500 nM capsaicin without 30 μ M TCO (ncAA) in media. **(A and B)** Neither TRPV1-T468tag-GFP (A) nor TRPV1-K464tag-GFP (B) generate Ca^{2+} responses in comparison to their transfection expression controls when supplemented with the pNEU plasmid but grown without TCO. All fluorescence experiment image sets are normalized to the brightness of their respective ionomycin image.



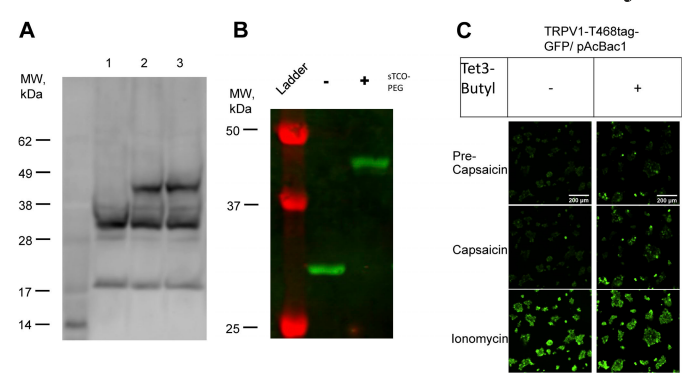


Figure S4. Western blots of gel-shift assay with sfGFP-N150tag/Tet3-Butyl and sTCO-PEG5K. (A) Extracts of sfGFP-N150tag/Tet3-Butyl expressed in HEK293T/17 pulse-labeled with sTCO-PEG5K. Lane 1 represents sfGFP-150tag expressed in the presence of NES-R284 (pAcBac1) and Tet3-Butyl ncAA. Lane 2 demonstrates that the addition of sTCO-PEG5k for 3 min induces a gel shift in the molecular weight band, and lane 3 shows the molecular weight shift of the sfGFP-N150tag after 10 min of incubation with sTCO-PEG5K. (B) Extraction of sfGFP-N150tag-FLAG/Tet3-Butyl with RIPA buffer and subsequent with (+) and without (-) sTCO-PEG5K labeling for 10 min before quenching with 1 mM Tet2 is shown after blotting for the C-terminal FLAG tag. (C) Control experiment of TRPV1-T468tag-GFP with pAcBac1 alone or supplemented with Tet3-Butyl in culture media. Ca²⁺ imaging of TRPV1 activity done as described in methods using Fluo4-AM and 500 nM capsaicin. All images are normalized to the brightness of ionomycin image. A single responding cell in the absence of Tet3-Butyl (-) was qualitatively deemed an outlier since cells otherwise showed no activity compared with Tet3-Butyl-treated cells (+), which showed robust activity with capsaicin. Source data are available for this figure: SourceData FS4.



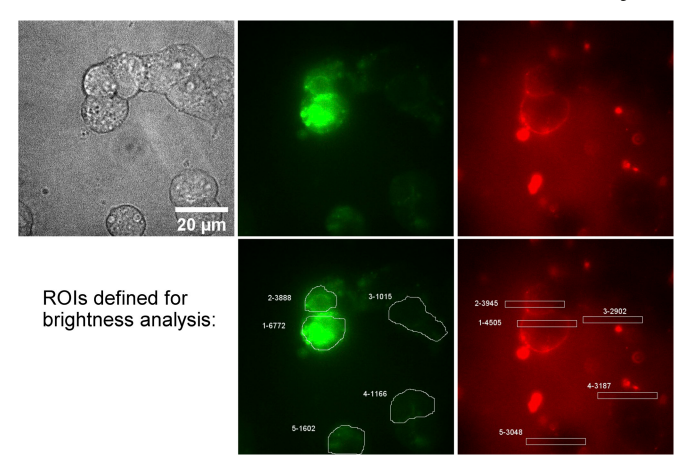


Figure S5. **Brightness correlation analysis of TRPV1-T468tag/Tet3-Bu-GFP expressing cells and labeled with sTCO-Cy5.** 100× objective images of HEK293T cells (DIC, top left) show varied degrees of GFP fluorescence from construct expression (GFP filter image, top middle) and correspondingly have different intensities of membrane labeling by sTCO-Cy5 (Cy5 filter image, top right). ROIs selected for mean GFP fluorescence in five cells (bottom middle) and Cy5 maximal pixel intensity (bottom right) are shown to illustrate our approach of capturing the full cell area to collect GFP fluorescence and selecting membrane regions of Cy5 that are representative of even, homogeneous dye distribution.



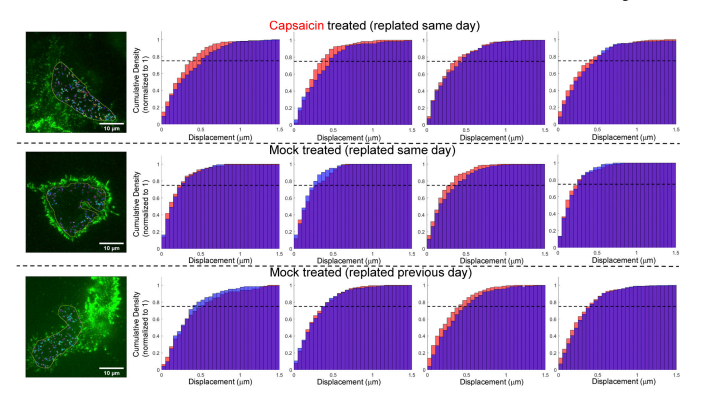


Figure S6. **Cumulative density histograms of track displacements before and after capsaicin treatment.** HEK293T cells expressing TRPV1exCellHalo are labeled with Alexa488 HaloTag ligand for single molecule tracking movies (see Materials and methods). 2.5-s movies (50 frames) are acquired, 15 s apart, before and after 5 μM capsaicin treatment to assess changes in TRPV1exCellHalo mobility. Cumulative density histograms (blue: pretreatment data; red: post-treatment data; purple: overlapping region in histograms) of track displacements are shown for three separate treatment conditions: (top row) capsaicin treatment with TRPV1exCellHalo expressing cells that are replated the same day onto poly-D-lysine coated coverslips; (middle row) mock treatment of TRPV1exCellHalo expressing cells replated onto coverslips 1 day earlier. A guideline at 0.75 is meant to help observe the upper-quartile values of the displacement distributions. Panels on the far left illustrate ROI-bounded tracks from cells that are paired with data in the blue histogram in the adjacent panel. Summary plots for these data are shown in Fig. 5 D.

Video 1. Image sequence capture under pre-capsaicin conditions in our capsaicin treatment sample. The first frame of the sequence is shown in Fig. 5 B and the analyzed track distances are displayed in the blue histogram of Fig. 5 C. Playback is at 0.5× speed.