Primary sensory neuron-specific interference of TRPV1 signaling by adeno-associated virus-encoded TRPV1 peptide aptamer attenuates neuropathic pain

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Abstract

Background: TRPV1 (transient receptor potential vanilloid subfamily member 1) is a pain signaling channel highly expressed in primary sensory neurons. Attempts for analgesia by systemic TRPV1 blockade produce undesirable side effects, such as hyperthermia and impaired heat pain sensation. One approach for TRPV1 analgesia is to target TRPV1 along the peripheral sensory pathway.

Results: For functional blockade of TRPV1 signaling, we constructed an adeno-associated virus (AAV) vector expressing a recombinant TRPV1 interfering peptide aptamer, derived from a 38mer tetrameric assembly domain (TAD), encompassing residues 735 to 772 of rat TRPV1, fused to the C-terminus of enhanced green fluorescent protein (EGFP). AAV-targeted sensory neurons expressing EGFP-TAD after vector injection into the dorsal root ganglia (DRG) revealed decreased inward calcium current and diminished intracellular calcium accumulation in response to capsaicin, compared to neurons of naïve or expressing EGFP alone. To examine the potential for treating neuropathic pain, AAV-EGFP-TAD was injected into fourth and fifth lumbar (L) DRGs of rats subjected to neuropathic pain by tibial nerve injury (TNI). Results showed that AAV-directed selective expression of EGFP-TAD in L4/L5 DRG neuron somata, and their peripheral and central axonal projections can limit TNI-induced neuropathic pain behavior, including hypersensitivity to heat and, to a less extent, mechanical stimulation.

Conclusion: Selective inhibition of TRPV1 activity in primary sensory neurons by DRG delivery of AAV-encoded analgesic interfering peptide aptamers is efficacious in attenuation of neuropathic pain. With further improvements of vector constructs and in vivo application, this approach might have the potential to develop as an alternative gene therapy strategy to treat chronic pain, especially heat hypersensitivity, without complications due to systemic TRPV1 blockade.

Keywords

Transient receptor potential vanilloid 1, neuropathic pain, dorsal root ganglion, primary sensory neurons, adeno-associated virus, gene therapy

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Introduction

Chronic pain represents a major health challenge. Large strides have been made in understanding mechanisms underlying neuropathic pain. This includes the identification of the TRPV1 (transient receptor potential vanilloid 1), a member of the vanilloid receptors that are highly expressed in primary sensory neuron somata and their fibers, as an important cellular integrative sensor for detecting noxious stimuli and transducing pain signals in the settings of various types of chronic pain. However, therapy by systemic TRPV1 blockade is accompanied by hyperthermia and impaired heat pain sensation, so an approach that can selectively block affected pain pathways is needed. Growing evidence shows that TRPV1 nociception in primary sensory neurons depends on its interaction with many other molecules, which together comprise the TRPV1 interaction, selectively reduce hypersensitive behavior in pain animal models, pharmacologically application of membrane-permeable iPAs (i.e., iPAs fused to cell-penetrating peptide) is hampered by their low stability and proteolytic lability in cells, and systemic peptide therapy cannot restrict the iPAs to the neural pathways responsible for transduction and transmission of nociceptive signals. As a result, potential off-target effects unrelated to pain can prevent sufficient dosing for effective antinociception, as in the case of hyperthermia from systemic TRPV1 blockade. Ideally, the therapeutic iPAs should be restricted in situ to the pathological sites, should be produced at stable concentrations that are sufficient to desensitize hyperactive pain signaling, and should have effects that are adequately sustained to avoid the need for frequently repeated injections. Gene therapy is a suitable strategy to achieve these outcomes, with recombinant adenovirus (AAV) demonstrating promise as one of the gene delivery tools for targeted applications.

Here, we combine a genetic strategy using AAV to transfer and persistently express a fluorescent TAD chimera (enhanced green fluorescent protein (EGFP)-TAD) selectively in primary sensory neurons, with an anatomic strategy for segmentally targeting neurons at the level of the painful condition via injection into the dorsal root ganglia (DRG), which has been shown to be a safe procedure in both rats and humans. We show that selective expression of TRPV1 TAD in lumbar (L) L4/L5 DRG neurons and their axonal projections provides effective and sustained attenuation of neuropathic pain induced by tibial nerve injury (TNI).

Materials and methods

Animals

Experiments were performed in adult male Sprague-Dawley rats (five–six weeks old; 125–150 g body weight) purchased from Charles River Laboratories (Wilmington, MA). All animal procedures were reviewed and approved by the Animal Care Committee of the Zablocki VA Medical Center Animal Studies Subcommittee and the Medical College of Wisconsin IACUC (Permission number: 3690-03). Rats were housed in standard 12-h cycle lighting and were allowed ad libitum access to food and water prior to and throughout the experimental protocol.

Construct and AAV production

To construct the AAV vector coding a chimeric EGFP-TAD expression cassette, we subcloned a DNA fragment encoding a 38mer peptide ranging the residues 735 to 772 of rat TRPV1 into BsrGI/SaI sites of a single-strand AAV expressing plasmid pAAV-CMV-EGFP (Cell Biolabs, San Diego, CA). This generated pAAV-CMV-EGFP-TAD that codes the EGFP-TAD fusion protein under the transcriptional control of the CMV promoter (Supplementary Figure S1(a)). The construct was sequenced to confirm the desired sequence and the correct reading frame. Plasmids pAAV-CMV-EGFP-TAD and pAAV-CMV-EGFP were used to package AAV2/6-EGFP-TAD and AAV2/6-EGFP as a control (subsequently referred to as AAV6-TAD and AAV6-EGFP, respectively) for in vivo injection. AAV vectors were produced and purified in our laboratory by previously described methods. This included AAV particle purification by optiprep ultracentrifugation and concentration by use of Centricon Plus-20 (Regenerated Cellulose 100,000 MWCO, Millipore, Billerica, MA).
AAV titer was determined by PicoGreen (life technologies, Carlsbad, CA) assay, and final aliquots were kept in 1 x phosphate buffered saline containing 5% sorbitol (Sigma-Aldrich, St. Louis, MO) and stored at –80°C. The titers of AAV6-TAD and AAV6-EGFP vectors were $1.04 \times 10^{13}$ GC/ml and $1.0 \times 10^{12}$ GC/ml, respectively. The same lot of viral preparation was used for all in vivo experiments. Vectors were evaluated for purity by sodium dodecyl sulfate polyacrylamide gel (SDS-PAGE) electrophoresis followed by silver stain using a Pierce silver stain kit (Fisher Scientific, Rockford, IL) according to manufacturer’s protocol. Comparable purity of AAV6-TAD and control AAV6-EGFP was confirmed by silver staining showing dominant viral Vp1, Vp2, and Vp3 proteins (95% for AAV6-TAD and 88% for AAV6-EGFP) (Supplementary Figure S1(b)). Infectivity was estimated by evaluating the percentage of EGFP-positive HEK293T cells 48 h after transduction of either AAV6-EGFP-TAD or AAV6-EGFP at a multiplicity of infection of 10,000 (Supplementary Figure S1(c)). By immunoblotting, the EGFP-TAD chimeric protein stable expression was confirmed in the lysates of HEK293T cells transduced with AAV6-EGFP-TAD with a mass that is $\sim 10$ kDa larger than EGFP (Supplementary Figure S1(d)).

**Microinjection of AAV vectors into DRGs**

Nearly all of the sciatic DRG perikarya resided in the L4 and L5 DRGs. AAV vector was microinjected into right L4 and L5 DRGs using previously described techniques. In brief, the surgically exposed intervertebral foramen was minimally enlarged by removal of laminar bone. The injection was performed through a micropipette that was advanced approximately 100 μm into the ganglion. Rats received L4 and L5 DRG injections of either AAV6-TAD or AAV6-EGFP (one vector per rat), consisting of 2 μl with adjusted titers containing a total of $2.0 \times 10^{10}$ GC particles. The injection was performed over a 5 min period using a Nanoliter 2000 microprocessor-controlled injector (World Precision Instruments, Sarasota, FL). Removal of the pipette was delayed for an additional 5 min to minimize extrusion of the injectate. Following the injection and closure of overlying muscle and skin, the animals were returned to the animal house where they remained for the experimental times. We have previously shown that AAV6 administered in this fashion produce transduction limited to the neurons of the injected DRGs, without significant glial transduction, likely due to high purity of vectors.

**Tibial nerve injury**

To model clinical traumatic painful peripheral neuropathy, we performed a TNI, an established model of peripheral nerve injury that is less severe than spared nerve injury. Animals were anesthetized using isoflurane at 4% induction and 2% maintenance. Under anesthesia, the right sciatic nerve was isolated under aseptic surgical conditions by blunt dissection of the femoral biceps muscle. The sciatic nerve and its three branches were isolated (sural, common peroneal, and tibial nerves), and only the tibial nerve was tightly ligated and transected distal to the ligation. The overlying muscle and skin were then sutured following surgery. Sham-operated rats were subjected to all preceding procedures without nerve ligation and transection.

**Pain behavioral evaluation**

Behavioral tests were performed in a blinded fashion. Animals were habituated in individual test compartments for at least 1 h before each testing. Stimuli were applied to the lateral margin of the plantar aspect of the foot in the sural area of innervation.

1. **Mechanical allodynia:** Mechanical withdrawal threshold testing (von Frey test) was performed using calibrated monofilaments (0.3, 0.5, 0.8, 1.0, 2.8, 5, 9, 14, and 24 g. Patterson Medical, Bolingbrook, Illinois). Animals were habituated in individual test compartments in an elevated mesh-bottomed platform with a grid for at least 1 h before testing. Beginning with the 2.8 g filament, filaments were applied to the plantar skin with just enough force to bend the fiber and held for 1 s. If a response was observed, the next smaller filament was applied, and if no response was observed, the next larger was applied, until a reversal occurred, defined as a withdrawal after a previous lack of withdrawal, or vice versa. Following a reversal event, four more stimulations were performed following the same pattern. The forces of the filaments before and after the reversal, and the four filaments applied following the reversal, were used to calculate the von Frey threshold. Rats not responding to any filament were assigned a score of 25 g.

2. **Mechanical hyperalgesia (Pin test):** Noxious punctate mechanical stimulation was performed using the point of a 22 g spinal anesthesia needle that was gently applied to the center of plantar surface of the hindpaw without penetrating the skin. Five applications were separated by at least 10 s, which was repeated after 2 min, making a total of 10 touches. For each application, the induced behavior was a very brisk, simple withdrawal with immediate return of the foot to the cage floor, or a sustained elevation with grooming that included licking and chewing, and possibly shaking, which lasted at least 1 s. This hyperalgesic behavior is specifically associated with place avoidance.
Hyperalgesia was quantified by tabulating hyperalgesia responses as a percentage of total touches.

3. Heat nociception (Hargreaves test): This was performed using a device designed for the purpose of identifying heat sensitivity (Paw Thermal Stimulator System, University Anesthesia Research & Development Group, San Diego, CA). Rats were placed on a temperature-regulated glass platform heated to 30°C, and the lateral plantar surface of hind paws stimulated with a radiant heat source (50 W halogen bulb) directed through an aperture. The time elapsed from initiation of the stimulus until withdrawal (withdrawal latency) as detected by a series of photocells was measured. Each hind paw was tested four times, and the withdrawal latency values averaged.

**Cutaneous capsaicin-induced behaviors**

An acute cutaneous inflammation was induced by hind-paw intradermal injection of 10 μg capsaicin (8-methyl-N-vanillyl-6-nonamide, Sigma-Aldrich), prepared in a solution of 7% Tween 80 and 93% saline as previously described.33 Animals were briefly anesthetized with isoflurane before injection to minimize their suffering. Capsaicin was injected into the plantar surface of the hindpaw in a volume of 10 μl. Immediately after injection, paw withdrawal responses to the 9 g von Frey monofilament in 10 times for a duration of 1 s with an inter-stimulus interval of approximately 1 s and to radiant heat applied on plantar skin were tested at 15, 30, 45, and 60 min. Paw withdrawal frequency (%) to von Frey stimulation was calculated as the numbers of paw withdrawals in 10 tests × 100. Only immediate, robust withdrawal responses from the von Frey stimulation were recorded as positive responses. Control experiments were done by vehicle injections using Tween 80 saline vehicle at the same volume as the capsaicin solution.

**Immunohistochemistry**

Immunohistochemistry staining was performed by a standard immunofluorescent labeling protocol, as previously described.34 Briefly, the ipsilateral and contralateral DRGs, the corresponding levels of the spinal cord, the sciatic nerve at middle thigh level, and hindpaw glabrous skin were removed and fixed in 10% neutral buffered formalin, embedded with paraffin, and processed into a series of 5 μm sections. Sections were deparaffinized and rehydrated, and antigen retrieval was achieved by microwave heating in 10 mM citrate buffer (pH 6.0). Following primary antibodies or IB4 were used for colocalization immunostaining: monoclonal GFP (1:400, Santa Cruz Biotechnology, SCB, Santa Cruz, CA), mouse monoclonal β3-tubulin (1:500, SCB), mouse monoclonal calcitonin gene-related peptide (1:600, SCB), rabbit polyclonal TRPV1 (1:500, ThermoFisher, Waltham, MA), or Alexa 594 conjugated isolectin GS-IB4 (IB4, 1 μg/ml, Life Technologies), with normal immunoglobulin G (IgG from same species as the first antibody) replacement of first antibody as the negative control. The appropriate fluorophore-conjugated (Alexa 488 or Alexa 594) secondary antibodies (Jackson Immunoresearch, West Grove, PA) were used to reveal the immune complex. The sections were examined, and images captured using a Nikon TE2000-S fluorescence microscope equipped with an Optronics Quantifire digital camera with filters suitable for selectively detecting the green, red, and blue fluorescence. For double-label colocalization, images from the same section but showing different antigen signals (different colors) were captured separately and overlaid. Colocalization was deemed for two proteins in areas stained with the combined color when the two images were overlaid.

**Western blotting**

Lysates of HEK293T cells and DRGs were extracted using 1× radioimmunoprecipitation assay (RIPA) buffer (20 mm Tris-HCl pH 7.4, 150 mm NaCl, 1% Nonidet P-40, 1% sodium deoxycholate, 0.1% SDS, with 0.1% Triton X100 and protease inhibitor cocktail). Protein concentration determined by using the BCA kit (Pierce, Rockford, IL). Western blotting of cell and DRG lysates (20 μg protein) was preceded by SDS-PAGE gel electrophoresis, transferred onto nitrocellulose, and probed with a polyclonal rabbit anti-GFP antibody (1:1,000; Cell Signaling, Danvers, MA) or rabbit polyclonal TRPV1 (1:500, ThermoFisher). Immunoreactive proteins were detected by enhanced chemiluminescence (Pierce, Rockford, IL) after incubation with horseradish peroxidase-conjugated secondary antibodies (1:2,000, SCB) and exposed to photographic film. Glyceraldehyde 3-phosphate dehydrogenase was used as a loading control. Densitometric analysis of TRPV1 protein levels in DRG homogenates was performed using ImageJ v1.46. All samples were normalized to glyceraldehyde 3-phosphate dehydrogenase (GAPDH) as control.

**Calcium imaging**

Capsaicin-evoked increases of intracellular calcium levels ([Ca2+]i) were compared in small to medium (25–40 μm in diameter) L4 and L5 DRG neurons dissociated from naïve rats and animals injected with AAV6-TAD or AAV6-EGFP. Dissociated neurons cultured in coverslips were loaded with Fura-2-AM (5 μM) and maintained in...
Tyrode’s solution containing (in mM): NaCl 140, KCl 4, CaCl2 2, glucose 10, MgCl2 2, and 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES) 10, with an osmolarity of 297 to 300 mOsm and pH 7.4. Recording was performed within 6 h of dissociation. Transduced neurons were identified by EGFP fluorescence. Capsaicin (50 nM and 100 nM) was applied locally (flow rate approximately 3/min) via a fine pipette positioned close to the cell being studied. The fluorophore was excited alternately with 340 nm and 380 nm wavelength illumination, and images were acquired at 510 nm using a cooled 12-bit digital at a rate of 3 Hz. The fluorescence ratio R for individual neurons was determined as the intensity of emission during 340 nm excitation (I340) divided by I380. and the Ca2+ concentration was estimated by as \( (K_d)(\beta)(R_{\text{min}})/(R_{\text{max}}-R) \), where \( \beta \) is the ratio of I380 at zero and saturating Ca2+ concentrations. Values of \( R_{\text{min}} \) / \( R_{\text{max}} \), and \( \beta \) were determined by in-situ calibration and were 0.38, 8.49 and 9.54, and Kd was 224 nm.35 Plasma membrane Ca2+-ATPase influence was eliminated by applying Tyrode’s with pH 8.8 during depolarization, while stable intracellular [Ca2+]i was maintained by simultaneously reducing bath Ca2+ concentration to 0.25 mM.

Voltage clamp of dissociated DRG neurons
After dissection in cold Ca2+/Mg2+-free HBBS (Life Technologies), ganglia were chopped, followed by incubating in 0.5 mg/ml liberase TM (Roche, Indianapolis, IN) in DMEM/F12 with glutaMAX (Life Technologies) for 30 min at 37°C, then with 1 mg/ml trypsin (Sigma-Aldrich) and 150 Kunitz units/ml DNase (Sigma-Aldrich) for another 10 min. After addition of trypsin inhibitor (Type II, Sigma-Aldrich), tissues were centrifuged, lightly triturated in neural basal media (1X) (Life Technologies) containing 2% (v:v) B27 supplement (50X) (Life Technologies), 0.5 mM glutamine (Sigma-Aldrich), 0.05 mg/ml gentamicin (Life technologies), and 10 ng/ml nerve growth factor 7S (Alomone Labs Ltd., Jerusalem, Israel). Cells were then plated onto poly-L-lysine (70–150 kDa, Sigma-Aldrich) coated coverslips and cultured at 37°C in 5% CO2. All neurons were studied 3 to 8 h after dissociation. Small to medium sensory neurons (25–40 μm in diameter) were used for this study. Electrodes with a resistance of 2 to 4 MΩ were pulled from borosilicate glass (Garner Glass Co., Claremont, CA) using a micropipette puller (P-97 Sutter Instrument Co; Novato, CA, USA) and fire polished. Recording was performed in the whole-cell configuration with an Axopatch 700B amplifier (Molecular Devices, Sunnyvale, CA). After whole-cell configuration was established, electrical compensation for the cell membrane capacitance and series resistance were initiated. Access resistance was typically between 5 and 7 MΩ and was 80% to 90% compensated. Since peak currents measured in our experiment were around 3 nA, voltage errors introduced by the residual uncompensated access resistance (<1.0 MΩ) were <3 mV and could not introduce major errors. Neurons with >10 MΩ access resistance after breakthrough were discarded. Experiments were performed 5 min after breakthrough, and at room temperature (~25°C). Signals were filtered at 2 kHz through a four-pole Bessel filter and digitized at 10 kHz with a Digidata 1440A A/D interface (Molecular Devices). Seals were achieved in modified Tyrode’s consisting of (in mM) NaCl 140, KCl 4, CaCl2 2, MgCl2 2, D-glucose 10, HEPES 10 at pH of 7.4, and with an osmolarity of 300 mOsm. Capsaicin (100 nM)-induced currents flowing through Ca2+ channels (ICa) were recorded using an extracellular solution consisting of (in mM) BaCl2 2, 4-amino-phenyridine 1, HEPES 10, tetrathylammonium chloride 140, HEPES 10 at pH of 7.4, and with an osmolarity of 300 mOsm. The internal pipette solution contained (in mM) CsCl 110, tetrathylammonium chloride 20, Mg-ATP 4, Na-GTP 0.3, ethylene glycolbis(2-aminoethylether)-N,N’,N’,N’-tetra-acetic acid 11, CaCl2 1, MgCl2 1, 4 Mg-ATP, 0.4 Li4-GTP, HEPES 10 at pH of 7.2, and with an osmolarity of 300 mOsm. Data from whole-cell ICa recordings were evaluated with Axograph X 1.3.5 (A xoGraph Scientific, Sydney, Australia), with which peak inward currents and charge transfer were measured. To correct for cell size, inward currents are expressed as a function of cell capacitance (pA/pF).

In vivo electrophysiological recordings of DH neurons
Rats were anesthetized with sodium pentobarbital (50 mg/kg, i.p.) with a catheter inserted into a femoral vein for supplemental administration of anesthetic doses (2 mg/kg, i.v.). Arterial blood pressure was monitored continuously using a PA-C10 transmitter (Data Sciences International (DSI), St. Paul, MN) inserted into a femoral artery. The trachea was cannulated through a midline cervical incision for ventilation with room air (Harvard 683 respirator), pneumothoraxes performed, and the rat was paralyzed with pancuronium bromide (0.1 mg/kg i.v., with supplemental doses of 0.01 mg/kg administered when spontaneous breathing was observed) prior to laminectomy. A surgical plane of anesthesia was reached prior to administration of pancuronium, as indicated by stability of blood pressure during stimulation. Supplemental doses of anesthetic were administered as indicate according to blood pressure changes. A heating pad was used to maintain body temperature at 37°C. For extracellular DH neuronal recordings, a laminectomy was performed from the T13 to the L3 vertebrae to expose the mid-lumbar
spinal cord, and a stabilizing stereotaxic clamp was applied to the spinal process rostral to the exposure. The dura was opened, and the cord was covered in warm mineral oil. First, activity evoked by mechanical stimulation of the receptive field was recorded. A single-barreled glass micropipette containing a carbon filament (7 µm diameter) for recording was advanced into the spinal cord using a microdrive, targeting lamina IV to VI of the L4 to L5 level at depths of 300 to 600 µm from the cord surface. Wide dynamic range neurons that responded to non-noxious and noxious stimulation in a graded manner were selected for study. Sequentially, the ipsilateral hindpaw was stimulated using cotton tip (stroking for 1 s at 0.5 Hz for 20 s), graded von Frey fibers (forces 0.07 g, 1.14 g, 8.0 g, and 60 g, each applied for 1 s at 0.5 Hz for 20 s), and venous clamp and arterial clamps that apply mild and moderate pinching force (each applied for 10 s one time). The various stimuli were applied following a 3 min rest interval. After recording evoked activity, spontaneous neuronal activity was recorded at two different longitudinal sites within the L4/5 recording area, and at four specified depths (300, 400, 500, and 600 µm from the cord surface) at each location, for 1 min at each location. This design was chosen to avoid sampling bias by going to these prespecified recording locations without reference to evocable activity.

Electrophysiological signals were directed to high impedance differential amplifiers (gain = 1000; 0.1–10 kHz passband), followed by filter/amplifiers (gain up to 400; high and low pass filtering 10 Hz–3 kHz). The amplifier output was displayed online and also directed to precision full-wave rectifiers and averaged using Bessel linear averaging filters (averaging interval = 100 ms) to obtain an online moving time average. Neuronal activity and the moving time average were continuously displayed/recorded along with arterial blood pressure with the CED Power 1401 mk II and Spike2 data acquisition system (Cambridge Electronic Design Limited, Cambridge, UK). Action potentials were isolated by setting the threshold above background noise, and individual units were identified by template matching using Spike2.

Statistical analysis

All statistical analysis was performed using Prism program (GraphPad Software, La Jolla, CA). Behavioral changes over time were analyzed by repeated measures parametric two-way analysis of variance (ANOVA) for von Frey and heat tests followed by Tukey’s post hoc test for multiple comparisons, and non-parametric Friedman’s ANOVA for pin test followed by Dunn’s post hoc test. For comparisons between groups, the effects of vector injection were characterized by area under the curve analysis. Specifically, for Early Treatment, measured behavioral values after the combined injury and vector injection were normalized to the values immediately preceding injury and injection. For Late Treatment (in which vector injection was performed 14 days after nerve injury), measured values after the vector injection were normalized to the values immediately before the injection. Calculated area under the curves were compared between vectors by Student’s t test for von Frey and heat and by Mann–Whitney U test for Pin. In electrophysiology experiments of DH neurons, baseline levels of neuronal spontaneous and evoked firing in rats were compared using Student’s t test. Statistical differences in the experiments of measurement of cytoplasmic Ca2+ concentration, whole-cell patch-clamp recording, and TRPV1 expression by immunoblots were analyzed by one-way ANOVA followed by Tukey’s post hoc test. Data were expressed as means ± SEM. A probability of p < 0.05 was considered as statistically significant.

Results

Construct design of AAV encoding TAD

TRPV1 is a tetrameric integral membrane protein, and its subunit organization and function are facilitated by interactions among C-terminal cytoplasmic domains,19,20,36–38 which contain the essential tetramerization sequence of TRPV1. Therefore, we aimed to block TRPV1 assembly by expressing TAD in the cytoplasm of DRG neurons. Two adjacent small peptide motifs mapped to TRPV1 cytosolic C-terminus (amino acids ranging 735–752 and 752–772, respectively), which are overlapped from each other by one amino acid, have been independently defined in prior reports as the critical TRPV1 TAD.16,19 Hence, we constructed an AAV encoding a 38mer TAD peptide (TRPV1 aa735–772), which was fused to EGFP in order to provide both a stable scaffold and a fluorescent tag for identifying neurons expressing EGFP-TAD driven under the cytomegalovirus (CMV) promoter. It is known that TRPV1 is highly expressed in various subtypes of primary sensory neurons, including C, Aδ, and some Aβ, and is distributed by axonal transport to both central presynaptic and peripheral terminals, although relatively limited TRPV1 expression in spinal cord interneurons and descending supraspinal fibers has recently been reported.39–41 AAV was packaged as serotype 6, because we have previously shown that after intraganglionic delivery, the AAV6 serotype provides efficient gene transfer to the full range of DRG neurons, including the nociceptive sub-populations and their axonal terminals, and highly purified AA6V with ubiquitous cell-potent CMV promoter leads to gene transfer predominantly to neurons without significant glial cell...
transduction in rat. Injection of AAV6-EGFP-TAD (subsequently referred to as AAV6-TAD) or control AAV6-EGFP into L4 and L5 DRGs of naïve rats did not alter thermal or mechanical sensitivity of the ipsilateral plantar skin during the four weeks postinjection observation period (Supplementary S2), nor did vector application appear to affect ambulation or limb posture, as evaluated by general observation.

**Sensory neuron expression of EGFP-TAD inhibits TRPV1 channel activity**

TRPV1 responds to capsaicin by increased nonselective Ca\(^{2+}\) permeability. We therefore examined the functional efficacy of EGFP-TAD expression in the primary sensory neurons by recording capsaicin-induced changes in intracellular calcium ([Ca\(^{2+}\)]\(_i\)) and whole-cell calcium current (I\(_{Ca}\)):

Small to medium sensory neurons (25–40 μm in diameter) were cultured from L4/L5 DRGs of naïve animals and from L4/L5 DRGs four weeks after they had been injected with either AAV6-TAD or AAV6-EGFP. Fura-2 imaging showed that transient increases in [Ca\(^{2+}\)]\(_i\) after stimulation with capsaicin (50 and 100 nM) were similar in naïve neurons and neurons expressing only EGFP (peak [Ca\(^{2+}\)]\(_i\) of 50 nM capsaicin: naïve neurons 1.02 ± 0.6 μM, neurons expressing EGFP 1.25 ± 0.2 μM, p > 0.05; peak [Ca\(^{2+}\)]\(_i\) of 100 nM capsaicin: naïve neurons 0.92 ± 0.3 μM, neurons expressing EGFP 0.95 ± 0.3 μM, p > 0.05) (Figure 1(a)). In contrast, neurons expressing EGFP-TAD showed a significantly decreased capsaicin-stimulated [Ca\(^{2+}\)]\(_i\) peak (0.46 ± 0.3 μM and 0.26 ± 0.3 μM for 50 and 100 nM of capsaicin, p < 0.001 and p < 0.01 vs. neurons expressing EGFP). Electrophysiological recording showed large inward capsaicin-induced currents in control neurons, and consistent with the [Ca\(^{2+}\)]\(_i\) data, peak inward current density was not different between neurons from naïve rats and neurons expressing EGFP ( naïve neurons: 8.52 ± 1.4 pA/pF, neurons expressing EGFP: 8.90 ± 1.73 pA/pF).

Figure 1. Expression of EGFP-TAD in sensory neurons inhibits TRPV1 channel activity and reduces response to plantar capsaicin. (a) and (b) Representative recording traces of intracellular calcium ([Ca\(^{2+}\)]\(_i\)) (a) and whole-cell calcium currents (I\(_{Ca}\)) (b) activated by 100 nM of capsaicin (Cap) in dissociated primary sensory neurons from naïve rat, and from AAV6-EGFP neurons and AAV6-TAD expressing neurons. Summarized data of Ca\(^{2+}\) transient exposed to 50 and 100 nM Cap (right panel of (a)) and whole-cell I\(_{Ca}\) response to 100 nM Cap (right panel of (b)) are shown as bar charts. Applications of Cap are shown in black bars. (c) and (d) Measurements of heat withdrawal latencies (c) and mechanical withdrawal frequency ((d), single 10 g von Frey filament) after hindpaw subcutaneous 10 μg Cap. Left panels are time course after injection of Cap, and right panels are summarized results at 15 min after injection. *p < 0.05, **p < 0.01, and ***p < 0.001.
and thermal stimuli. To further test the efficacy of TRPV1 and sensitizes the skin to noxious mechanical stimulation is reduced in sensory neurons expressing EGFP-TAD.

Intradermal administration of capsaicin activates TRPV1 and sensitizes the skin to noxious mechanical and thermal stimuli. To further test the efficacy of AAV6-TAD, we examined heat and mechanical responses after intraplantar capsaicin (10 μg) in naïve animals and those injected with AAV6-EGFP or AAV6-TAD (28 days after injection, except for two rats at 29 days for AAV-GFP and one rat 26 days after AAV-TAD; five animals per group). The time latency to withdraw the paw from radiant heat projected on the plantar skin was markedly reduced after intradermal capsaicin in naïve animals, and this was unchanged in rats with AAV6-EGFP injected in the ipsilateral L4 and L5 DRGs. However, this hypersensitivity to heat was diminished in rats injected with AAV6-TAD (Figure 1(c)). To test the effect of plantar capsaicin on mechanical sensitivity, the rate of withdrawal response to plantar stimulation with a 9 g von Frey filament was measured. Capsaicin increased the response frequency in naïve and AAV6-EGFP injected rats, but this mechanical hypersensitivity was diminished in AAV6-TAD injected rats (Figure 1(d)). Together, these results indicate that AAV-directed selective expression EGFP-TAD in DRG neurons leads to desensitization of TRPV1 neurons to capsaicin and suggest an analgesic potential of TAD against both heat and mechanical hypersensitivity.

Neuropathic pain is alleviated by AAV-encoded TAD expression

To test the hypothesis that AAV-mediated TAD expression selectively in the primary sensory neurons can prevent neuropathic pain following peripheral nerve injury, rats were injected with either AAV6-TAD or AAV6-EGFP (2.0 × 10^10 viral particles per DRG) at the same time as performing TNI (referred hereafter to this design as Early Treatment). Nearly all of the sciatic DRG perikarya reside in the L4 and L5 DRGs, so these two DRGs were injected. Sensitivity to heat and mechanical cutaneous stimulation was evaluated prior to TNI and DRG injection and was followed for eight weeks after injection of the vectors. Results showed that, after TNI surgery, all animals injected with AAV6-EGFP developed elevated sensitivity to heat, mechanical allodynia to innocuous mechanical stimulation (von Frey), and hyperalgesia to noxious mechanical stimulation (Pin), which lasted the full period of observation (Figure 2(a)). Animals injected with AAV6-TAD also exhibited hypersensitivity to heat stimulation and mechanical allodynia and hyperalgesia after TNI. However, hypersensitivity to these stimuli was significantly attenuated compared to the animals injected with the AAV6-EGFP control vector (Figure 2(a)). These findings suggest that application of AAV6-TAD early in the development of injury-induced neuropathic pain limits the development of hypersensitivity to heat and mechanical stimulation.

We next evaluated the ability of AAV-mediated TAD expression to reverse established neuropathic pain (termed Late Treatment). In this design, rats received intragangliionic vector injection (L4/L5 DRGs) 14 days after TNI, followed by seven weeks of sensory behavior evaluation. Results showed that there was no significant effect on mechanical hyperalgesia (Pin; Figure 2(b)), and that the average degree of hyperalgesia (%) for the final four determinations of hyperalgesia (days 35 through 56 after TNI) was less for the early treatment (30 ± 4%) than for late treatment (40% ± 7%; p = 0.029), which indicates that early treatment is important for mechanical hyperalgesia. The attenuating effect of Late Treatment on mechanical allodynia (von Frey) was significant (von Frey, Figure 2(b)), but allodynia during the final four determinations was nonetheless greater after Late Treatment 4.0 ± 0.4 g than when AAV6-TAD was delivered early (7.4 ± 0.7 g; p = 0.0006). For heat hypersensitivity, Late Treatment with AAV6-TAD substantially reversed latency (Figure 2(b)), and the average degree of latencies for the final four determinations in Late Treatment were comparable (9.5 ± 0.1 s) to the average latencies in Early Treatment (9.7 ± 0.05 s; p = 0.0488), indicating that AAV6-TAD retains analgesic efficacy against hypersensitivity to heat even in established neuropathic pain.

AAV6-TAD produces long-lasting transgene expression

Immunohistochemical characterization on L4/L5 DRGs, lumbar spinal cord, and hindpaw glabrous skin from rats eight weeks after TNI plus Early Treatment showed that AAV6-TAD efficiently and persistently transduced DRG neurons that include sub-populations positive for the TRPV1, the small peptidergic neuron marker calcitonin gene-related peptide, and the nonpeptidergic small neuron marker isolectin B4 (IB4), as well as large-sized neurons immunolabeled by NF-200 (Figure 3(a) to (e)). In neurons of all size groups, TRPV1 expression was found in neurons that were highly co-labeled for EGFP-TAD in sections from the TNI group, suggesting that TRPV1 is expressed not only in the non-myelinated and thinly myelinated C- and Aδ-fibers but also in the myelinated Aβ-fibers following nerve injury. This may in part explain the observed effectiveness of AAV6-TAD treatment on mechanical hypersensitivity in TNI animals. The in vivo transduction rate for...
AAV6-TAD at eight-week after TNI followed by vector injection was 51% ± 8% of total neuronal profiles (positive for β3-tubulin) within sections showing the entire ganglion (n = 3 DRGs, five sections per DRG). Examination of the corresponding spinal cord revealed that EGFP-TAD fibers terminate predominantly in the superficial laminae, some in deeper laminae, as well as dorsal column, of ipsilateral dorsal horn (DH) (Figure 3(f)), whereas no EGFP-TAD signal was observed in DH neurons. Abundant EGFP-TAD was also detected in the skin sections of ipsilateral hindpaws (Figure 3(g)), but no transgene product was detected in brain and liver (Data not shown). These findings validate stable expression of the transgene product (EGFP-TAD) restricted in the peripheral sensory nervous pathway ipsilateral to injection. Similar patterns of EGFP expression in the DRG neurons and their axonal projections were detected following AAV6-EGFP injection.
Western blots of DRG tissue homogenates prepared from rats eight weeks after TNI plus vector injection (Early Treatment) identified the EGFP-TAD fusion protein based on the size disparity between EGFP-TAD and EGFP, indicating stability of EGFP-TAD protein in vivo during eight weeks of expression (Figure 4).
TRPV1 protein level in TNI-DRGs injected with AAV6-TAD was not different from levels in TNI-DRGs injected with AAV6-EGFP or in sham control DRGs (Figure 4, n = 6 per group). However, TRPV1 protein level in TNI-DRGs (L5) injected with AAV6-EGFP was significantly elevated compared with control DRGs. To verify this TRPV1 elevation in TNI DRG was not due to vector injection, immunoblotting and immunohistochemistry analyses were performed in the ipsilateral L5 DRGs harvested four weeks after TNI alone. After injury, TRPV1 immunopositivity appeared in all neuronal size groups, in contrast to control DRG section in which TRPV1 expression was mainly limited to small-sized neurons (Figure 5(a) to (c)). The relative TRPV1 protein level was 59% ± 23% higher after TNI compared to control DRGs (n = 6 per group, p < 0.01; Figure 5(d)). It has previously been reported that spinal nerve ligation upregulates TRPV1 heat function in injured IB4-positive DRG neurons, and that both TRPV1 mRNA and protein are upregulated in L5 DRGs ipsilateral to spared nerve injury. TNI produces DRGs with co-mingled injured and uninjured axons, so our showing elevated TRPV1 level following TNI are compatible with prior findings and support the importance of TRPV1 in sensitization and pain transmission following TNI nerve injury.

Effect of AAV6-TAD on DH neuronal activity

Pain signaling is supported by TRPV1 not only by initiating activity in peripheral sensory neuron terminals but also by aiding neurotransmission in the spinal cord DH. TRPV1 is enriched in the central presynaptic terminals of the DH, where it facilitates the transmission of noxious stimuli, and nerve injury-induced TRPV1 receptor upregulation occurs at central presynaptic sites of primary afferents leading to an enhancement of excitatory signaling in the spinal cord. Prior studies of animals with neuropathic pain have observed abnormal activity in neurons of the lamina V area of the DH, including elevated spontaneous activity and evoked activity following cutaneous stimulation. We therefore examined firing properties of deep DH neurons in TNI rats after Late Treatment with AAV6-TAD. Extracellular recordings of evoked and spontaneous activity of spinal cord DH neurons were obtained from anesthetized TNI rats at a timepoint nine weeks after TNI and seven weeks after injection of either AAV6-EGFP or AAV6-TAD. Total spontaneous activity was recorded ipsilateral to the TNI at two different longitudinal locations within the L4/L5 levels of the spinal cord and within the laminae IV to VI at four prechosen depths from the surface at each longitudinal location. This design was used in order to avoid sampling bias towards active areas. The activity of 51 such sites from TNI rats treated by AAV6-TAD (n = 8) and 55 sites from the TNI rats by AAV6-EGFP injection (n = 7) were recorded. This showed decreased spontaneous activity in AAV6-TAD-treated TNI animals (3.2 ± 0.1 spikes/s) compared to recordings from AAV6-EGFP-treated TNI animals (3.8 ± 0.2 spikes/s, p < 0.05; Figure 6(a)). In the same animals, activity evoked by receptive field mechanical stimuli (brush, graded von Frey filaments, and pinch by two graded vascular clamps) was used to identify single units as wide dynamic range neurons. Comparison of these evoked responses between animals injected with
AAV6-TAD versus AAV6-EGFP, however, did not reveal significant differences (Figure 6(b)).

**Discussion**

Our general aim is to develop primary sensory neuron-specific AAV-encoded iPA treatment for neuropathic pain, for which TRPV1 was chosen as a target because it is a critical signaling node for pain transduction and transmission in primary sensory neurons. Additionally, this genetic approach offers a unique opportunity to test the role of DRG neurons in neuropathic pain. The results presented here validate that AAV6-delivered, sensory neuron-specific expression of a fluorescent TAD, consisting of a 38mer iPA designed to interrupt TRPV1 subunit assembly, can reduce TRPV1 activity in primary sensory neurons. Furthermore, single injection of AAV6-TAD into lumbar DRGs leads to long-term TAD expression and sustained attenuation of neuropathic hypersensitivity to both heat and mechanical stimulation. A detailed mechanistic analysis remains to be documented, but analgesic effects of AAV-encoded TAD in DRG neurons may be mediated by disrupting TRPV1 function at multiple sites along the entire neuron, including decreased ectopic activity in the soma, modulated impulse generation at the peripheral terminals, and inhibited presynaptic neurotransmission in the DH.

Overall, our findings add further evidence that support primary sensory neuron TRPV1 as an important target for pain management. Other genetic-based and ligand-based TRPV1 silencing/ablating strategies have been proposed for targeting sensory neurons, such as TRPV1 RNA interference and combined application of resiniferatoxin and lidocaine derivative QX314 treatment, which efficiently provide relief of pathological pain. However, RNA interference gene knockdown results in posttranscriptional silencing of all TRPV1 functions, and QX314 can produce cytotoxicity. In contrast, the AAV-iPA strategy may provide a more finely tuned regulation of neuronal function because it blocks only the selected protein–protein interaction, while leaving other roles of this multifunctional protein unaffected, such as TRPV1’s neuroprotective role.

TRPV1 not only encodes noxious heat in primary sensory neurons but also responds to chemicals that trigger pain after neuroinflammation and additionally...
is implicated in neuropathic pain. Our results show that targeting DRG TRPV1 signaling specifically by TAD is effective for attenuating heat hypersensitivity and also can diminish mechanical hypersensitivity. Convergent evidence suggests that TRPV1 may function as a molecular integrator for multiple types of sensory inputs including mechanotransduction, although this is not yet resolved. Previous reports show that intraganglionic resiniferatoxin induces analgesia for both thermal and mechanical stimulation, while intrathecal AAV-shRNA targeting TRPV1 produces heat analgesia but mechanical pain sensation remained intact. Pain alleviation by targeting sensory neurons with TAD interference may be mechanistically different from these various approaches. Specifically, it has been shown that the C-terminus of TRPV1 contains overlapping segments that are necessary for interaction with multiple molecules that are involved in pain pathology by various mechanisms. Therefore, there should be interacting effects of signal transduction components by our 38mer TAD motif. These may involve, for example, the putative binding sites for calmodulin, A-kinase anchoring proteins, and lipid mediators such as endovanilloids and lysophosphatidic acid, as well as phosphorylation sites. Furthermore, the TAD domain of TRPV1 is highly conserved with the corresponding C-terminal amino acid sequence of other TRPV family members, especially TRPV4 that has 63% residue identities with TRPV1-TAD (Figure 7), although other transient receptor potential members lack this sequence homology. Thus, the functional relevance of AAV-mediated TAD interference might extend beyond TRPV1 channel. TRPV4 is known to mediate mechanotransduction and plays a crucial role in a painful peripheral neuropathy. TRPV4 is reported highly colocalized with TRPV1 in DRG neurons, and biochemical and biophysical evidence also indicate a close proximity between TRPV4 and TRPV1 in DRG neurons. Therefore, a detailed examination of the ion channels and receptors involved in the pathophysiology of pain conditions related to the TRPV1 interactome is a suitable high priority for future investigation. It was not our goal to identify the subgroup of neurons that account for the therapeutic efficacy of our AAV ipA approach. Future research can employ AAV serotypes with tropism for distinct neuronal populations and selective promoters to isolate the one or more specific neuronal populations that need to be treated. Additionally, as there is 30% to 60% homology for TRPV2, TRPV3, and TRPV4 at the TAD, compared to TRPV1, we cannot attribute the entire analgesic efficacy of EGFP-TAD to action only on TRPV1.

Our preliminary electrophysiological observations show that AAV6-TAD reduces spontaneous activity of DH neurons in the deep laminae of the nucleus proprius. This may reflect an underlying therapeutic mechanism contributing to the analgesic effect of AAV6-TAD since spontaneous activity of these neurons is a feature of various neuropathic pain conditions. Spontaneous activity in DH neurons could be triggered by activity originating in peripheral sensory fibers following injury. We did not identify an effect of AAV6-TAD on firing evoked by graded mechanical stimuli, although this negative finding could result from the inherently heterogeneity of sensory units and resulting high variance in the data. Additionally, these recordings were performed at a late timepoint when the effect of AAV6-TAD treatment was decreasing, and in the Late Treatment experimental design. Both of these features diminished the efficacy of AAV-TAD treatment shown in behavioral evaluations, so electrophysiological differences from the AAV-EGFP-treated controls would be expected to be limited in these animals. A greater effect
of AAV6-TAD on DH neuron activity might have been seen in Early Treatment animals, and at earlier time-points. Nonetheless, the observation of diminished spontaneous activity despite these limitations may indicate limiting such activity may be a critical feature of AAV6-TAD analgesia.

We observed analgesic efficacy in two experimental designs that reflect possible clinical application. Administering AAV6-TAD at the time of the injury could be employed clinically when nerve injury is an unavoidable result of surgery, such as limb amputation. Administering AAV6-TAD after a neuropathic pain state is established represents the more typical clinical setting. Although we saw an incomplete analgesic effect of AAV6-TAD, improved efficacy could be achieved by increasing transduction efficiency with repeated injections. Generally, a gene therapy approach based on anatomically targeted AAV delivery of iPAs for controlling specific protein–protein interactions in sensory neurons has high potential for translation into a clinically useful modality in treating chronic pain. AAV has a very favorable safety profile, and there is a growing experience of clinical application using this vector. The high specificity of iPAs should limit off-target effects. Intraganglionic injections are a minimally invasive procedure that is routinely performed by appropriately trained physicians. The fact that the widely used procedure of foraminal injection (also known as selective spinal nerve block) commonly results in intraganglionic injection but nerve damage is rare attests to the safety of injecting within the DRG. Ultimately, the broad range of known protein–protein interactions involved in pain signaling may allow substantial control of sensory neuron pain signaling by AAV-delivered iPAs.

Author’s Contributions
HY and QHH conceived and designed the experiments. HY, HX, ZL, FW, GF, CJR, DC, and QHH conducted the experimental investigation and developed methodology. HY, HX, GF, BP, and QHH analyzed the data. Funding acquisition by QHH and HY. HY, QHH, and CJS wrote the paper. HX and ZL contributed equally to this work.

Declaration of Conflicting Interests
The author(s) declared no potential conflicts of interest with respect to the research, authorship, and/or publication of this article.

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Supplementary Materials

Figure S1. Design and preparation of AAV vectors. (a) Schematic outline of AAV vectors. The vector genomes of AAV-TAD and AAV-EGFP are flanked by AAV2 inverted terminal repeats (ITR). Transgene cassettes encode EGFP-TAD separated by a 22aa linker between EGFP and TAD or EGFP-linker alone downstream of a chimeric intron enhancing transcription driven by the cytomegalovirus (CMV) promoter, and contain polyadenylation signal element from the human growth hormone (GHpA). The amino acid sequence of a 38mer TAD (red) is shown on the top and the linker sequence (back) in the middle of two vectors. (b) Silver stain result of purified AAV6-TAD and AAV6-EGFP. This revealed 3 virion protein bands of Vp1, Vp2, and Vp3, with molecular weight of 87, 72 and 62 KDa, respectively. (c) GFP expression by infectivity estimation upon transduction to HEK293T cells of either AAV6-EGFP or AAV6-TAD at a multiplicity of infection of 10,000. (d) Western analysis on lysates of HEK293T cells transduced by vectors show EGFP immunoreactivity at distinct molecular weights (MWs) for expressed EGFP (lane 1) versus EGFP-TAD (lane 2 and 3). Arrows point to the expected size bands for EGFP-TAD and EGFP (top panel), and glyceraldehyde-3-phosphate dehydrogenase (GAPDH, bottom panel) as a loading control.
Figure S2. Baseline sensory evaluation after vector injection into DRGs. (a–c) Sensory sensitivity evaluated by Heat, von Frey, and Pin tests after vector injection in non-injured rats. (a) Hargreaves test for heat sensitivity, (b) vF test for innocuous mechanical sensitivity, and (c) Pin test for the evaluation of noxious mechanical sensitivity before and 28-d after vector injection, compared to sham-operated (no injection) animals (n = 5 rats per group).

Figure S3. EGFP expression following intraganglionic injection with control vector AAV6-EGFP. Representative IHC images show EGFP expression (a, green) in DRG neurons (b, merged with phase-contrast image) and lumbar spinal cord (c, white matter of spinal cord was pseudocolored to blue) after 8 weeks of intraganglionic injection with AAV6-EGFP. Scale bars: 200 μm.